P/ NT COOPERATION TREAT

From the INTERNATIONAL BUREAU

PCT

NOTIFICATION OF ELECTION

(PCT Rule 61.2)

To:

Commissioner US Department of Commerce United States Patent and Trademark Office, PCT

2011 South Clark Place Room

CP2/5C24

Arlington, VA 22202

ETATS-UNIS D'AMERIQUE

in its capacity as elected Office

Date of mailing (day/month/year)

03 November 2000 (03.11.00)

International application No. PCT/EP00/02441

International filing date (day/month/year) 20 March 2000 (20.03.00)

Applicant's or agent's file reference

D 1321 PCT

Priority date (day/month/year) 19 March 1999 (19.03.99)

Applicant

DE VEYLDER, Lieven et al

nder

The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland

Authorized officer

Claudio Borton

Telephone No.: (41-22) 338.83.38

Facsimile No.: (41-22) 740.14.35



(PCT Article 18 and Rules 43 and 44)

Applicant's or agent's file reference D 1321 PCT	FOR FURTHER see Notification of Transmittal of International Search Report (Form PCT/ISA/220) as well as, where applicable, item 5 below.					
International application No.	International filing date (day/month/year)	(Earliest) Priority Date (day/month/year)				
PCT/EP 00/02441						
Applicant						
CROPDESIGN N.V.		·				
This International Search Report has been	n prepared by this International Searching Auth	nority and is transmitted to the applicant				
according to Article 18. A copy is being tra	ansmitted to the International Bureau.	•				
This International Search Report consists	of a total of 4 sheets.					
1 000	a copy of each prior art document cited in this	report.				
Basis of the report						
	international search was carried out on the bas ess otherwise indicated under this item.	sis of the international application in the				
the international search w Authority (Rule 23.1(b)).	as carried out on the basis of a translation of the	he international application furnished to this				
b. With regard to any nucleotide an was carried out on the basis of the	d/or amino acid sequence disclosed in the in e sequence listing:	ternational application, the international search				
	nal application in written form.					
filed together with the inte	rnational application in computer readable form	n.				
furnished subsequently to	this Authority in written form.					
	this Authority in computer readble form.					
international application a	sequently furnished written sequence listing do s filed has been furnished.	•				
the statement that the info	ormation recorded in computer readable form is	s identical to the written sequence listing has been				
2. Certain claims were fou	nd unsearchable (See Box I).					
3. Unity of invention is lac	king (see Box II).					
4. With regard to the title,						
the text is approved as su	bmitted by the applicant.					
the text has been establis	hed by this Authority to read as follows:					
		·				
5. With regard to the abstract,						
X the text is approved as su	bmitted by the applicant.					
the text has been established, according to Rule 38.2(b), by this Authority as it appears in Box III. The applicant may, within one month from the date of mailing of this international search report, submit comments to this Authority.						
6. The figure of the drawings to be publ		7				
as suggested by the appli	cant.	None of the figures.				
X because the applicant fail	ed to suggest a figure.					
because this figure better	characterizes the invention.					

P P 00/02441

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C12N15/29 C12N15/82

C12N9/12

C12N5/10

A01H5/00

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) IPC 7 C12N A01H

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

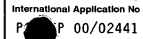
EPO-Internal, BIOSIS

C. DOCUMENTS CONSIDERED TO BE RELEVANT				
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.		
X	WO 97 02354 A (MEDICAL RES COUNCIL; LATHANGUE NICHOLAS BARRIE (GB)) 23 January 1997 (1997-01-23) the whole document	28-36		
A	WO 98 41642 A (VEYLDER LIEVEN DE ;VLAAMS INTERUNIV INST BIOTECH (BE); INZE DIRK () 24 September 1998 (1998-09-24) cited in the application see the whole document; esp. pp.8-16	1-45		
A	<pre>WO 99 13083 A (VEYLDER LIEVEN DE ;INZE DIRK (BE); CROPDESIGN N V (BE); SEGERS GER) 18 March 1999 (1999-03-18) the whole document</pre>	1–45		
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X Further documents are listed in the continuation of box C.	Patent family members are listed in annex.		
 Special categories of cited documents: "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed 	 "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family 		
Date of the actual completion of the international search 5 September 2000	Date of mailing of the international search report 21/09/2000		
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Kania, T		

International	Application N
PQ	00/02441

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	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
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Α	WO 98 03631 A (SALK INST FOR BIOLOGICAL STUDI) 29 January 1998 (1998-01-29) the whole document	1-45
Α	√ WO 92 09685 A (UNIV AUSTRALIAN) 11 June 1992 (1992-06-11) cited in the application the whole document	1-45
A	RIOU-KHAMLICHI C ET AL: "Cytokinin activation of Arabidopsis cell division through a D-type cyclin" SCIENCE, US, AMERICAN ASSOCIATION FOR THE ADVANCEMENT OF SCIENCE,, vol. 283, no. 5407, 5 March 1999 (1999-03-05), pages 1541-1544, XP002134537 ISSN: 0036-8075 cited in the application the whole document	1-45
Α	FRANCIS DENNIS: "Regulation of the cell cycle plant development" BIOSIS, ACTA PHARMACEUTICA, vol. 45, no. 2, 1995, XP002140190 abstract	1-45
A	DOONAN J: "PLANT GROWTH: ROOTS IN THE CELL CYCLE" CURRENT BIOLOGY,GB,CURRENT SCIENCE,, vol. 6, no. 7, 1 July 1996 (1996-07-01), pages 788-789, XP002045511 ISSN: 0960-9822 the whole document	1-45
A	SEGERS GERDA ET AL: "The Arabidopsis cyclin-dependent kinase gene cdc2bAt is preferentially expressed during S and G-2 phases of the cell cycle" PLANT JOURNAL,GB,BLACKWELL SCIENTIFIC PUBLICATIONS, OXFORD, vol. 10, no. 4, 1996, pages 601-612, XP002138663 ISSN: 0960-7412 cited in the application the whole document	1-45
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A	GENSCHIK PASCAL ET AL: "Cell cycle-dependent proteolysis in plants: Identification of the destruction box pathway and metaphase arrest produced by the proteasome inhibitor MG132." PLANT CELL, vol. 10, no. 12, December 1998 (1998-12), pages 2063-2075, XP002146545 ISSN: 1040-4651 the whole document	1-45
Α	FOWLER M R ET AL: "The plant cell cycle in context." MOLECULAR BIOTECHNOLOGY, vol. 10, no. 2, October 1998 (1998-10), pages 123-153, XP000939061 ISSN: 1073-6085 see esp. parts 10-12	1-45
P,X	WO 99 58681 A (RAMIREZ PARRA ELENA ;XIE QI (ES); CONSEJO SUPERIOR INVESTIGACION () 18 November 1999 (1999-11-18) cited in the application see the whole document; esp. p.5 1.26 - p.6 1.2; claims 1-4	1,2,13, 19-45

Info on on patent family members

EP 00/02441

Patent document cited in search repo		Publication date		Patent family member(s)	Publication date
WO 9702354	Α	23-01-1997	AÚ CA EP JP	6237396 A 2224940 A 0835319 A 11509089 T	05-02-1997 23-01-1997 15-04-1998 17-08-1999
W0 9841642	Α	24-09-1998	AU EP	6730198 A 0972060 A	12-10-1998 19-01-2000
WO 9913083	Α	18-03-1999	AU EP	9537498 A 1007684 A	29-03-1999 14-06-2000
WO 9803631	Α	29-01-1998	AU BR CA EP	3960597 A 9710872 A 2260287 A 0929663 A	10-02-1998 17-08-1999 29-01-1998 21-07-1999
W0 9209685	A	11-06-1992	AU AU CA EP JP US US	657722 B 9046291 A 2097286 A 0559729 A 6504430 T 5750862 A 6087175 A	23-03-1995 25-06-1992 30-05-1992 15-09-1993 26-05-1994 12-05-1998 11-07-2000
W0 9958681	Α	18-11-1999	AU	3828099 A	29-11-1999

From the INTERNATIONAL SEARCHING AUTHORITY	PCT			
To: VOSSIUS & PARTNER Siebertstrasse 4 81675 München GERMANY EINGEGAN Vossius & Par 2 1. Sep. 200	oer On the Declaration			
Frist 21.11, bearb.: 21.10. Je	Date of mailing (day/month/year) 21/09/2000			
Applicant's or agent's file reference	21/0//2000			
D 1321 PCT	FOR FURTHER ACTION See paragraphs 1 and 4 below			
International application No. PCT/EP 00/02441	International filing date (day/month/year) 20/03/2000			
Applicant				
CROPDESIGN N.V.				
1. X The applicant is hereby notified that the International Search Report has been established and is transmitted herewith. Filling of amendments and statement under Article 19: The applicant is entitled, if he so wishes, to amend the claims of the International Application (see Rule 46): When? The time limit for filing such amendments is normally 2 months from the date of transmittal of the International Search Report; however, for more details, see the notes on the accompanying sheet. Where? Directly to the International Bureau of WIPO 34. chemin des Colombettes 1211 Geneva 20, Switzerland Fascimile No.: (41–22) 740.14.35 For more detailed Instructions, see the notes on the accompanying sheet. 2. The applicant is hereby notified that no International Search Report will be established and that the declaration under Article 17(2)(a) to that effect is transmitted herewith. 3. With regard to the protest against payment of (an) additional fee(s) under Rule 40.2, the applicant is notified that the protest together with the decision thereon has been transmitted to the International Bureau together with the applicant's request to forward the texts of both the protest and the decision thereon to the designated Offices. In o decision has been made yet on the protest; the applicant will be notified as soon as a decision is made.				
Shortly after 18 months from the priority date, the international application will be published by the International Bureau. If the applicant wishes to avoid or postpone publication, a notice of withdrawal of the international application, or of the priority claim, must reach the International Bureau as provided in Rules 90bis.1 and 90bis.3, respectively, before the completion of the technical preparations for international publication. Within 19 months from the priority date, a demand for international preliminary examination must be filed if the applicant				
wishes to postpone the entry into the national phase until 30 months from the priority date (in some Offices even later). Within 20 months from the priority date, the applicant must perform the prescribed acts for entry into the national phase before all designated Offices which have not been elected in the demand or in a later election within 19 months from the priority date or could not be elected because they are not bound by Chapter II.				
Nam and mailing address of the International Searching Authority European Patent Office. P.B. 5818 Patentlaan 2 NL-2280 HV Rijswijk Tel. (+31-70) 340-2040. Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Andria Overbeeke-Siepkes			

NOTES TO FORM PCT/ISA/220

These Notes are intended to give the basic instructions concerning the filing of amendments under article 19. The Notes are based on the requirements of the Patent Cooperation Treaty, the Regulations and the Administrative Instructions under that Treaty. In case of discrepancy between these Notes and those requirements, the latter are applicable. For more detailed information, see also the PCT Applicant's Guide, a publication of WIPO.

In these Notes, "Article", "Rule", and "Section" refer to the provisions of the PCT, the PCT Regulations and the PCT Administrative Instructions respectively.

INSTRUCTIONS CONCERNING AMENDMENTS UNDER ARTICLE 19

The applicant has, after having received the international search report, one opportunity to amend the claims of the international application. It should however be emphasized that, since all parts of the international application (claims, description and drawings) may be amended during the international preliminary examination procedure, there is usually no need to file amendments of the claims under Article 19 except where, e.g. the applicant wants the latter to be published for the purposes of provisional protection or has another reason for amending the claims before international policiation. Furthermore, it should be emphasized that provisional protection is available in some States only.

What parts of the international application may be amended?

Under Article 19, only the claims may be amended.

During the international phase, the claims may also be amended (or further amended) under Article 34 before the International Preliminary Examining Authority. The description and drawings may only be amended under Article 34 before the International Examining Authority.

Upon entry into the national phase, all parts of the international application may be amended under Article 28 or, where applicable, Article 41.

When?

Within 2 months from the date of transmittal of the international search report or 16 months from the priority date, whichever time limit expires later. It should be noted, however, that the amendments will be considered as having been received on time if they are received by the International Bureau after the expiration of the applicable time limit but before the completion of the technical preparations for international publication (Rule 46.1).

Where not to file the amendments?

The amendments may only be filed with the International Bureau and not with the receiving Office or the International Searching Authority (Rule 46.2).

Where a demand for international preliminary examination has been its filed, see below.

How?

Either by cancelling one or more entire claims, by adding one or more new claims or by amending the text of one or more of the claims as filed.

A replacement sheet must be submitted for each sheet of the claims which, on account of an amendment or amendments, differs from the sheet originally filed.

All the claims appearing on a replacement sheet must be numbered in Arabic numerals. Where a claim is cancelled, no renumbering of the other claims is required. In all cases where claims are renumbered, they must be renumbered consecutively (Administrative Instructions, Section 205(b)).

The amendments must be made in the language in which the international application is to be published.

What documents must/may accompany the amendments?

Letter (Section 205(b)):

The amendments must be submitted with a letter.

The letter will not be published with the international application and the amended claims. It should not be confused with the "Statement under Article 19(1)" (see below, under "Statement under Article 19(1)").

The letter must be in English or French, at the choice of the applicant. How ver, if the language of the international application is English, the letter must be in English; if the language of the international application is French, the letter must be in French.

Notes to Form PCT/ISA/220 (first sheet) (January 1994)

The letter must indicate the differences between the claims as filed and the claims as amended. It must, in particular, indicate, in connection with each claim appearing in the international application (it being understood that identical indications concerning several claims may be grouped), whether

- (i) the claim is unchanged;
- (ii) the claim is cancelled;
- (iii) the claim is new;
- (iv) the claim replaces one or more claims as filed;
- (v) the claim is the result of the division of a claim as filed.

The following examples illustrate the manner in which amendments must be explained in the accompanying letter:

- [Where originally there were 48 claims and after amendment of some claims there are 51]:
 "Claims 1 to 29, 31, 32, 34, 35, 37 to 48 replaced by amended claims bearing the same numbers; claims 30, 33 and 36 unchanged; new claims 49 to 51 added."
- [Where originally there were 15 claims and after amendment of all claims there are 11]:
 "Claims 1 to 15 replaced by amended claims 1 to 11."
- [Where originally there were 14 claims and the amendments consist in cancelling some claims and in adding new claims]:
 - "Claims 1 to 6 and 14 unchanged; claims 7 to 13 cancelled; new claims 15, 16 and 17 added." or "Claims 7 to 13 cancelled; new claims 15, 16 and 17 added; all other claims unchanged."
- 4. [Where various kinds of amendments are made]: "Claims 1-10 unchanged; claims 11 to 13, 18 and 19 cancelled; claims 14, 15 and 16 replaced by amended claim 14; claim 17 subdivided into amended claims 15, 16 and 17; new claims 20 and 21 added."

"Statement under article 19(1)" (Rule 46.4)

The amendments may be accompanied by a statement explaining the amendments and indicating any impact that such amendments might have on the description and the drawings (which cannot be amended under Article 19(1)).

The statement will be published with the international application and the amended claims.

it must be in the language in which the international appplication is to be published.

It must be brief, not exceeding 500 words if in English or if translated into English.

It should not be confused with and does not replace the letter indicating the differences between the claims as filed and as amended. It must be filed on a separate sheet and must be identified as such by a heading, preferably by using the words "Statement under Article 19(1)."

It may not contain any disparaging comments on the international search report or the relevance of citations contained in that report. Reference to citations, relevant to a given claim, contained in the international search report may be made only in connection with an amendment of that claim.

Consequence if a demand for international proliminary examination has already been filed

If, at the time of filing any amendments under Article 19, a demand for international preliminary examination has already been submitted, the applicant must preferably, at the same time of filing the amendments with the International Bureau, also file a copy of such amendments with the International Preliminary Examining Authority (see Rule 52.2(a), first sentence).

Consequence with regard to translation of the international application for entry into the national phase

The applicant's attention is drawn to the fact that, where upon entry into the national phase, a translation of the claims as amended under Article 19 may have to be furnished to the designated/elected Offices, instead of, or in addition to, the translation of the claims as filed.

For further details on the requirements of each designated/elected Office, see Volume II of the PCT Applicant's Guide.

PCT

INTERNATIONAL SEARCH REPORT

(PCT Article 18 and Rules 43 and 44)

Applicant's or agent's file reference D 1321 PCT	FOR FURTHER see Notification of Transmittal of International Search Report (Form PCT/ISA/220) as well as, where applicable, item 5 below.		
International application No.	International filing date (day/mont	h/year) (Earliest) P	riority Date (day/month/year)
PCT/EP 00/02441	20/03/2000		19/03/1999
Applicant CROPDESIGN N.V.			
This International Search Report has be according to Article 18. A copy is being to This International Search Report consist	ransmitted to the International Burea	eets.	ansmitted to the applicant
Basis of the report			
a. With regard to the language, the	e international search was carried out nless otherwise indicated under this i	on the basis of the interreem.	national application in the
the international search Authority (Rule 23.1(b)).	was carried out on the basis of a tran	slation of the internationa	al application furnished to this
b. With regard to any nucleotide a was carried out on the basis of the contained in the international furnished subsequently the statement that the suinternational application	nd/or amino acid sequence disclosine sequence listing: ional application in written form. ernational application in computer reso this Authority in written form. o this Authority in computer readble to absequently furnished written sequences filed has been furnished. formation recorded in computer read-	adable form. orm. ce listing does not go bey	.· yond the disclosure in the
	und unsearchable (See Box I).		
3. Unity of invention is la	cking (see Box II).		
4. With regard to the title,			
the text is approved as s	ubmitted by the applicant.		
the text has been establi	shed by this Authority to read as folk	ws:	
the text has been establi	ubmitted by the applicant. shed, according to Rule 38.2(b), by to date of mailing of this international blished with the abstract is Figure No	search report, submit cor	s in Box III. The applicant may, mments to this Authority.
as suggested by the app			None of the figures.
because the applicant fa	iled to suggest a figure.		
	. Character at invention.		

EP 00/02441

A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 C12N15/29 C12N15/82

C12N9/12

C12N5/10

A01H5/00

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) IPC 7 C12N A01H

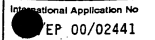
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Date of the actual completion of the international search 5 September 2000	Date of mailing of the international search report 21/09/2000
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2	Authorized officer
NL 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Kania, T



C.(Continua	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	EP UC	
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International Application No /EP 00/02441

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n on patent family members

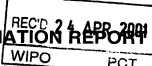
International Application No
/EP 00/02441

Patent document cited in search rep		Publication date		Patent family member(s)	Publication date
WO 9702354	A	23-01-1997	AU CA EP JP	6237396 A 2224940 A 0835319 A 11509089 T	05-02-1997 23-01-1997 15-04-1998 17-08-1999
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WO 9913083	· A	18-03-1999	AU EP	9537498 A 1007684 A	29-03-1999 14-06-2000
WO 9803631	A	29-01-1998	AU BR CA EP	3960597 A 9710872 A 2260287 A 0929663 A	10-02-1998 17-08-1999 29-01-1998 21-07-1999
WO 9209685	A	11-06-1992	AU AU CA EP JP US US	657722 B 9046291 A 2097286 A 0559729 A 6504430 T 5750862 A 6087175 A	23-03-1995 25-06-1992 30-05-1992 15-09-1993 26-05-1994 12-05-1998 11-07-2000
WO 9958681	Α	18-11-1999	AU	3828099 A	29-11-1999

TENT COOPERATION TRE

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INTERNATIONAL PRELIMINARY EXAMINATION REPORT



(PCT Article 36 and Rule 70)

Applicant's or agen	nt's file reference			Sac Natifier	ation of Transmittal of International				
D 1321 PCT	•	FOR FURTHER AC	CTION		Examination Report (Form PCT/IPEA/416)				
International applica	ation No.	International filing date (day/month/	'year)	Priority date (day/month/year)				
PCT/EP00/024	41	20/03/2000			19/03/1999				
International Patent Classification (IPC) or national classification and IPC C12N15/82									
Applicant									
CROPDESIGN	N.V.								
This international preliminary examination report has been prepared by this International Preliminary Examining Authority and is transmitted to the applicant according to Article 36.									
2. This REPOR	RT consists of a total of	8 sheets, including this	cover sh	eet.					
 This report is also accompanied by ANNEXES, i.e. sheets of the description, claims and/or drawings which have been amended and are the basis for this report and/or sheets containing rectifications made before this Authority (see Rule 70.16 and Section 607 of the Administrative Instructions under the PCT). These annexes consist of a total of sheets. 									
3. This report of	ontains indications relat	ting to the following iten	ns:						
ı 🛭 E	Basis of the report								
. II 🛛 F	Priority								
111 🗆 1	Non-establishment of or	pinion with regard to no	velty, inve	entive step a	and industrial applicability				
ıv □ ı	Lack of unity of inventio	n							
	Reasoned statement un citations and explanatio			ovelty, inve	ntive step or industrial applicability;				
VI ⊠ (Certain documents cite	d			Ĭ.				
VII 🗆 (Certain defects in the in	ternational application							
VIII ⊠ (Certain observations on	the international applic	ation						
Date of submission	of the demand		Date of c	ompletion of t	his report				

20.04.2001 02/10/2000 Name and mailing address of the international Authorized officer preliminary examining authority: European Patent Office D-80298 Munich Roscoe, R Tel. +49 89 2399 - 0 Tx: 523656 epmu d Fax: +49 89 2399 - 4465 Telephone No. +49 89 2399 2554

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No. PCT/EP00/02441

I. Basi	s ft	h re	p rt
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1.	the and	With regard to the elements of the international application (Replacement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report since they do not contain amendments (Rules 70.16 and 70.17)): Description, pages:						
	1-6	8	as originally filed					
	Cla	ims, No.:						
	1-4	5	as originally filed					
	Dra	wings, sheets:						
	1/5-	5/5	as originally filed					
2.			juage, all the elements marked above were available or furnished to this Authority in the international application was filed, unless otherwise indicated under this item.					
	These elements were available or furnished to this Authority in the following language: , which is:							
	☐ the language of a translation furnished for the purposes of the international search (under Rule 23.1(b)).							
		the language of pu	ublication of the international application (under Rule 48.3(b)).					
		the language of a 55.2 and/or 55.3).	translation furnished for the purposes of international preliminary examination (under Rule					
3.	With regard to any nucleotide and/or amino acid sequence disclosed in the international application, the international preliminary examination was carried out on the basis of the sequence listing:							
		contained in the in	ternational application in written form.					
		filed together with	the international application in computer readable form.					
		furnished subsequ	ently to this Authority in written form.					
		furnished subsequ	ently to this Authority in computer readable form.					
		☐ The statement that the subsequently furnished written sequence listing does not go beyond the disclosure in the international application as filed has been furnished.						
		The statement tha listing has been fu	t the information recorded in computer readable form is identical to the written sequence rnished.					
4.	The	amendments have	resulted in the cancellation of:					
		the description,	pages:					
		the claims,	Nos.:					

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No. PCT/EP00/02441

		the drawings,	sheets:					
5. This report has been established as if (some of) the amendments had not been made, since they have considered to go beyond the disclosure as filed (Rule 70.2(c)):								
		(Any replacement shareport.)	eet contai	ning such	amendments must be referred to under item 1 and annexed to this			
6.	Add	litional observations, if	necessar	y:				
II.	Pric	ority						
1.		This report has been prescribed time limit t			priority had been claimed due to the failure to furnish within the			
		☐ copy of the earlie	er applicat	ion whose	e priority has been claimed.			
		☐ translation of the	earlier ap	plication	whose priority has been claimed.			
2.		This report has been been found invalid.	establishe	ed as if no	priority had been claimed due to the fact that the priority claim has			
	Thu date		his report,	the inter	national filing date indicated above is considered to be the relevant			
3.		Additional observations, if necessary: see separate sheet						
٧.		Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; sitations and explanations supporting such statement						
1.	Stat	ement						
	Nov	relty (N)	Yes: No:		9, 11, 12, 14, 16-18, 45-68 1-8, 10, 13, 15, 19-44			
	Inve	entive step (IS)	Yes: No:		18, 45-68 1-17, 19-44			
	Indu	ustrial applicability (IA)	Yes: No:	Claims Claims	1-68			
2.	Cita	tions and explanations	S					

VI. Certain documents cited

see separate sheet

1. Certain published documents (Rule 70.10)

International application No. PCT/EP00/02441

and / or

2. Non-written disclosures (Rule 70.9)

see separate sheet

VIII. Certain observations on the international application

The following observations on the clarity of the claims, description, and drawings or on the question whether the claims are fully supported by the description, are made: see separate sheet

I. Basis

The documents mentioned in the present International Preliminary Examination Report are numbered as in the search report, i.e. D1 corresponds to the first document of the search report etc.

II. Priority

Cannot be acknowledged for claims 9-18 (and to those parts of further claims dependent on or referring back to claims 9-18) unless applicant can show where the priority document discloses these specific CDKs and cyclins and the specific combinations thereof.

V. Reasoned statement on Novelty, Inventive Step and Industrial Applicability

- Novelty (Art.33(2) PCT)

D1 discloses complexes of p53 with the transcription factors DP-1 or E2F-1 (and other related proteins). Use complexes to assay for modulators of these interactions. These transcription factors are important for regulation of cell cycle progression. Constructs expressing both transcription modulator (p53) and transcription factor (e.g. DP1) are disclosed on p.5-6 (e.g.). Further disclosed are DNA constructs, host cells, methods for preparing complex,... D1 anticipates claims 28-30, 32-36, 44.

D2 discloses use of 2-hybrid assay to find new products interacting with Cdc2aAt (a CDK). Thus, the cells in which assay gives positive result express cell cycle interacting proteins from separate nucleic acids. Found novel plant CKS homolog in Arabidopsis (CKS1At). Transgenic plants were constructed overexpressing plant-specific cyclin dependent kinase (e.g. Cdc2b) or overexpressing CKS1At (see p.11). On p.12, it is stated that in a preferred embodiment, one or more cell cycle genes or plant cell cycle genes are specifically used to modulate endoreduplication in plants (see also claim 19). D2 anticipates claims 1-5, 10, 20-27, 29-34, 36.



INTERNATIONAL PRELIMINARY **EXAMINATION REPORT - SEPARATE SHEET**

International application No. PCT/EP00/02441

D3 discloses the same 2-hybrid assay as D2. Identified th65 which binds and is phosphorylated by CDK. As D2, suggests expressing one or more DNA sequences encoding cell cycle regulatory proteins functional in plants. Cited against same claims as D2.

D5 discloses controlling plant growth by overexpressing p34cdc2a (which found to be limiting for cell division). States that invention extends to other cell cycle control proteins including p13suc1 (p34 complexes with this yeast protein), cyclin, cdc25 or combinations thereof separately or together with p34. Bottom p.9 suggests transformation of monocot protoplasts with a p34-like molecule and perhaps if necessary a cyclin-encoding gene. D5 anticipates claims 1-8, 15, 20-27, 31, 34, 37-44.

D13 is presently relevant only to those claims not entitled to priority from 19.03.99. A 2-hybrid system is disclosed wherein plant E2F interacts with plant Rb. Top of p.6 suggests that plant E2F may be modified alone or in combination with modification of levels of Rb (see also claim 4). D13 anticipates claims 13 and 19-45 (insofar as these claims are dependent on or refer back to claim 13).

Inventive Step (Art.33(3) PCT)

The prior art already suggests coexpression of multiple cell cycle interacting proteins to promote plant growth (see e.g. D2, D3), yet does not explicitly suggest coexpression of both partners of a cdc/cyclin complex. Insofar as use type A CDK and cyclin partner, inventive step could be acknowledged based on the unexpected effect observed. Type A CDK levels were not necessarily expected to be rate-limiting, and thus coexpression thereof with cyclin (cyclins are generally considered rate-limiting) surprisingly enhances growth. The claims are however virtually all much far broader than this inventive subject-matter. With regard to coexpression of other cell cycle interacting complex formers, no surprising effect has been established and hence, in view of the prior art no inventive step can be acknowledged. Specifically, type B CDK levels would be expected to be ratelimiting (as are cyclin levels in general) - hence obvious to attempt cooverexpression of type B CDK with Cyclin partner(s). Hence, of those claims which are novel, the following claims are considered to encompass non-inventive

subject-matter: 9, 11, 12, 14, 16 and 17

All claims relating to coexpression constructs or mixtures of constructs expressing CDK and cyclin partners will not considered inventive even if minor modifications can overcome the novelty problems. Coexpression of 2 members of a complex is obvious basis for study thereof (see e.g. D1)

Industrial Applicability (Art.33(4) PCT)

The present claims appear to have industrial applicability.

Certain documents VI.

In accordance with Rule 70.10, PCT, applicants attention is drawn to the following document(s):

WO-A-99/58681 (Publication date, 18.11.99; Priority date, 08.05.98 & 11.05.98; Filing date, 07.05.99)

VIII. Certain observations

Clarity (Art.6 PCT)

Claims 1-4 lack the essential feature of the invention i.e. CDK & cyclin overexpression. This feature is considered essential since it is surprising that coexpression leads to an improved result (basis for acknowledging inventive step). However, this surprising effect was only shown in CDK/cyclin context and not in vague context of cell-cycle interacting proteins. If it is truly surprising then it is not a basis for extrapolation to other proteins.

Claims 28-30, 32-45 lack the essential feature of the application, i.e. the combination of expression of CDK and cyclin to achieve an unexpected increase in e.g. growth. Many individual constructs have already been used in the prior art and unspecified expression constructs comprising known genes are not inventive anyway.

Claim 36 - product defined by process of production. However, the complexes were already known prior to application and thus not novel. Products cannot be rendered novel via their process of production.

Support in Description (Art.6 PCT)

The only actual data on enhanced plant growth provided by applicant relates to CycB1 and Cdc2aAt co-overexpression. Applicant claims that the enhanced growth achieved is surprising for 2 reasons: (i) since Cdc2a constitutively expressed, not expected to be rate-limiting, (ii) Doerner et al., 1996 showed CycB1 expression to be critical (this was not found under applicants experimental conditions). In spite of unexplained inconsistencies with previous data (ii), it is conceded that CycB1 and Cdc2aAt overexpression does have a surprising beneficial effect. The claims are however broad in scope and it is not considered that applicants at best meagre data supports such a breadth of protection.

There are a large number of cell cycle interacting proteins, many of which interact to form transient or stable complexes (see e.g. review article D12). Expression patterns of these proteins vary widely insofar as they have been established at all. Even amongst different CDK types, totally different patterns of expression are observed. Although Cdc2a is constitutively expressed, this is not the case for Cdc2bAt, which is expressed during limited interval of cell cycle (see D9). Similarly, in rice, Cdc2Os1 is constitutive and Cdc2Os2 cell cycle specific (D10). Hence, no evidence is provided that will get the same effect here. Extension of claimed subject-matter beyond A-type CDK-Cyclin interactions is thus not supported.

PATENT COOPERATION TREATY

From the INTERNATIONAL PRELIMINARY EXAMINING AUTHORITY

To:

VOSSIUS & PARTNEF Siebertstr. 4 D-81765 München ALLEMAGNE EINGEGANGEN Vossius & Partner

2 3. April 2001

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NOTIFICATION OF TRANSMITTAL OF THE INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Rule 71.1)

Date of malling

(day/month/year)

20.04.2001

Applicant's or agent's file reference

D 1321 PCT

IMPORTANT NOTIFICATION

International application No. PCT/EP00/02441

International filing date (day/month/year) 20/03/2000

Priority date (day/month/year)

19/03/1999

Applicant

CROPDESIGN N.V.

- 1. The applicant is hereby notified that this International Preliminary Examining Authority transmits herewith the international preliminary examination report and its annexes, if any, established on the international application.
- 2. A copy of the report and its annexes, if any, is being transmitted to the International Bureau for communication to all the elected Offices.
- 3. Where required by any of the elected Offices, the International Bureau will prepare an English translation of the report (but not of any annexes) and will transmit such translation to those Offices.

4. REMINDER

The applicant must enter the national phase before each elected Office by performing certain acts (filing translations and paying national fees) within 30 months from the priority date (or later in some Offices) (Article 39(1)) (see also the reminder sent by the International Bureau with Form PCT/IB/301).

Where a translation of the international application must be furnished to an elected Office, that translation must contain a translation of any annexes to the international preliminary examination report. It is the applicant's responsibility to prepare and furnish such translation directly to each elected Office concerned.

For further details on the applicable time limits and requirements of the elected Offices, see Volume II of the PCT Applicant's Guide.

Name and mailing address of the IPEA/

Authorized officer

Emslander, S

European Patent Office D-80298 Munich

Tel. +49 89 2399 - 0 Tx: 523656 epmu d

Fax: +49 89 2399 - 4465

Tel.+49 89 2399-8718

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INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

International application No. PCT/EP00/02441 International Patent Classification (IPC) or national classification and IPC C12N15/82 Applicant CROPDESIGN N.V. 1. This international preliminary examination report has been prepared by this International Preliminary Examinit and is transmitted to the applicant according to Article 36. 2. This REPORT consists of a total of 8 sheets, including this cover sheet. This report is also accompanied by ANNEXES, i.e. sheets of the description, claims and/or drawings white been amended and are the basis for this report and/or sheets containing rectifications made before this A (see Rule 70.16 and Section 607 of the Administrative Instructions under the PCT). These annexes consist of a total of sheets. 3. This report contains indications relating to the following items:		fication of Transmittal of International ary Examination Report (Form PCT/IPI		FOR FURTHER	Applicant's or agent's file reference D 1321 PCT			
PCT/EP00/02441		Priority date (day/month/year)	date (day/month/year)	International filing da	eation No			
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Name and mailing address of the international preliminary examining authority: European Patent Office	STATE A SOURS MAZE		Authorized officer	nal	ining authority:	exami		
D-80298 Munich Tel. +49 89 2399 - 0 Tx: 523656 epmu d Fax: +49 89 2399 - 4465	O STANDARD STANDARD	(!	Roscoe, R	56 epmu d)298 Munich +49 89 2399 - 0 Tx; 52365	D-80 Tel.	<u>)</u>	



I. Basis	of the	report
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1.	the . and	With regard to the elements of the international application (Replacement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report since they do not contain amendments (Rules 70.16 and 70.17)): Description, pages:						
	1-68	3	as originally filed					
	Clai	ms, No.:						
	1-45	5	as originally filed					
	Dra	wings, sheets:						
	1/5-	5/5	as originally filed					
2.			guage, all the elements mar international application wa					
	These elements were available or furnished to this Authority in the following language: , which is:							
		the language of a	translation furnished for the	purposes of the inte	rnational search (un	der Rule 23.1(b)).		
		the language of p	ublication of the internationa	al application (under F	Rule 48.3(b)).			
		the language of a 55.2 and/or 55.3)	translation furnished for the	purposes of internat	ional preliminary exa	amination (under Rule		
3.			cleotide and/or amino acid ry examination was carried			application, the		
		contained in the i	nternational application in w	ritten form.				
		filed together with	the international application	n in computer readab	le form.			
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		furnished subseq	uently to this Authority in co	mputer readable form	ı.			
			at the subsequently furnishe application as filed has beer		sting does not go be	yond the disclosure in		
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EXAMINATION REPORT

		the drawings,	sheets:		•			
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5.					ome of) the amendments has filed (Rule 70.2(c)):	ad not been mad	e, since they have bee	
		(Any replacement she report.)	eet contain	ning such	amendments must be refe	rred to under iten	n 1 and annexed to this	
6.	Ado	litional observations, if	necessar	/:			·	
H.	Pric	ority			· .			
1.		This report has been prescribed time limit t			priority had been claimed	due to the failure	to furnish within the	
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	Thu dat	• •	this report,	the interi	national filing date indicate	d above is consid	lered to be the relevant	
3.		ditional observations, i e separate sheet	f necessar	y:			·	
۷.		asoned statement un ations and explanatio			ith regard to novelty, inv th statement	entive step or in	dustrial applicability;	
1.	Sta	tement						
	No	velty (N)	Yes: No:	Claims Claims	9, 11, 12, 14, 16-18, 45-6 1-8, 10, 13, 15, 19-44	38		
	Inv	entive step (IS)	Yes: No:	Claims Claims	18, 45-68 1-17, 19-44	•		
		lustrial applicability (IA) Yes: No:	Claims Claims	1-68			
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2.		ations and explanatior e separate sheet	is .					

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VI.



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D13 is presently relevant only to those claims not entitled to priority from 19.03.99. A 2-hybrid system is disclosed wherein plant E2F interacts with plant Rb. Top of p.6 suggests that plant E2F may be modified alone or in combination with modification of levels of Rb (see also claim 4). D13 anticipates claims 13 and 19-45 (insofar as these claims are dependent on or refer back to claim 13).

Inventive Step (Art.33(3) PCT)

The prior art already suggests coexpression of multiple cell cycle interacting proteins to promote plant growth (see e.g. D2, D3), yet does not explicitly suggest coexpression of both partners of a cdc/cyclin complex. Insofar as use type A CDK and cyclin partner, inventive step could be acknowledged based on the unexpected effect observed. Type A CDK levels were not necessarily expected to be rate-limiting, and thus coexpression thereof with cyclin (cyclins are generally considered rate-limiting) surprisingly enhances growth. The claims are however virtually all much far broader than this inventive subject-matter. With regard to coexpression of other cell cycle interacting complex formers, no surprising effect has been established and hence, in view of the prior art no inventive step can be acknowledged. Specifically, type B CDK levels would be expected to be ratelimiting (as are cyclin levels in general) - hence obvious to attempt cooverexpression of type B CDK with Cyclin partner(s). Hence, of those claims which are novel, the following claims are considered to encompass non-inventive

Claim 36 - product defined by process of production. However, the complexes were already known prior to application and thus not novel. Products cannot be rendered novel via their process of production.

Support in Description (Art.6 PCT)

The only actual data on enhanced plant growth provided by applicant relates to CycB1 and Cdc2aAt co-overexpression. Applicant claims that the enhanced growth achieved is surprising for 2 reasons: (i) since Cdc2a constitutively expressed, not expected to be rate-limiting, (ii) Doerner et al., 1996 showed CycB1 expression to be critical (this was not found under applicants experimental conditions). In spite of unexplained inconsistencies with previous data (ii), it is conceded that CycB1 and Cdc2aAt overexpression does have a surprising beneficial effect. The claims are however broad in scope and it is not considered that applicants at best meagre data supports such a breadth of protection.

There are a large number of cell cycle interacting proteins, many of which interact to form transient or stable complexes (see e.g. review article D12). Expression patterns of these proteins vary widely insofar as they have been established at all. Even amongst different CDK types, totally different patterns of expression are observed. Although Cdc2a is constitutively expressed, this is not the case for Cdc2bAt, which is expressed during limited interval of cell cycle (see D9). Similarly, in rice, Cdc2Os1 is constitutive and Cdc2Os2 cell cycle specific (D10). Hence, no evidence is provided that will get the same effect here. Extension of claimed subject-matter beyond A-type CDK-Cyclin interactions is thus not supported.

PCT





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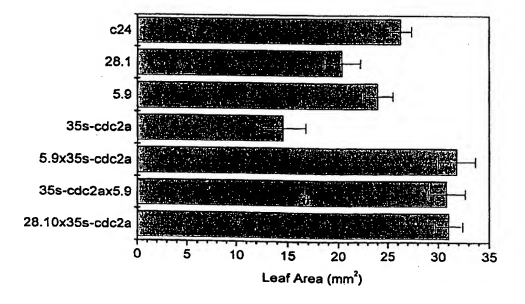
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(57) Abstract

Described is a method for promoting or modifying plant growth and/or yield and/or architecture in plants comprising the increased expression of at least two cell cycle interacting proteins, in particular of a protein kinase, e.g. CDK, and a cyclin. Transgenic plants are provided obtainable by this method and displaying increased cell division rates and growth rates. In addition, harvestable parts and propagation material of the above-mentioned plant as well as the use of the provided cells, tissues and plants for the production of biomass, secondary metabolites or additives for plant culturing in plant culture.

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Method for enhancing and/or improving plant growth and/or yield or modifying plant architecture

Field of the invention

The current invention relates to a method for modifying, preferably for enhancing or promoting plant growth and/or yield in plants and for modifying their architecture and to the transgenic plants obtainable by this method.

The invention concerns the simultaneous ectopic expression and/or overexpression of at least two cell cycle interacting proteins capable of forming a complex and specifically a CDK and an interacting cyclin; said co-overexpression results in an unexpected growth and architectural characteristics such as enhanced root and/or shoot growth in plants.

Several documents are cited throughout the text of this specification. Each of the documents cited herein (including any manufacturer's specifications, instructions, etc.) are hereby incorporated by reference; however, there is no admission that any document cited is indeed prior art as to the present invention.

Background of the invention

Cell cycle - General

Higher plants are multicellular organisms, hence their growth is a function of the rate at which new cells are produced. There is no doubt that the regulation of the cell division cycle plays a crucial role in plant growth regulation. WO 00/56905 PCT/EP00/02441

2

The eukaryotic cell division cycle can be divided in four main phases: the G1 phase or first gap phase, the S phase during which the DNA is duplicated, the G2 phase or second gap phase, and the M phase during which karyo- and cytokinesis take place. The major checkpoints regulating the progression through the cell cycle are situated at the G1/S and G2/M transitions. If the conditions are inadequate for the cell to continue its cycle, a block can occur at one or both transition points. The cell cycle can also be blocked at other transition points which had not until recently been considered important, namely M/G1. Passage through the G1/S and G2/M boundaries is dependent on the activity of cyclin dependent kinases (CDKs). A prerequisite for CDK activity, which by itself is regulated by phosphorylation and dephosphorylation events, is binding to a regulatory subunit, a cyclin. Generally cyclins oscillate at both transcriptional and protein level in a cell cycle phase dependent manner. The association of a cyclin with a CDK not only confers activity but also contributes to the substrate specificity of the CDK complex and its subcellular localisation.

CDKs

Intensive cloning efforts have identified a large number of CDK proteins in diverse plant species, among which at least five types can be distinguished on the basis of their sequences (for a compilation see Segers et al., 1997). In the model plant, *Arabidopsis thaliana*, two CDKs, each belonging to a different family, have been characterised. One such example is the *CDC2a*At gene, which contains the conserved PSTAIRE amino acid motif, and is constitutively expressed during the cell cycle at transcriptional and protein level. However, the associated kinase activity is maximal at the G1/S and G2/M transitions, suggesting a role at both checkpoints (Hemerly *et al.*, 1993; Burssens et al., 1998; Segers *et al.*; 1996). *CDC2b*At contains a PPTALRE motif and its mRNA levels are preferentially present during S and G2 phase (Segers *et al.*, 1996 and references cited therein). The protein follows the transcriptional level but the CDC2bAt kinase activity becomes only maximal during mitosis, implying a role during the M phase.

3

Cyclins

Numerous cDNAs encoding putative cyclins have been found in a diverse range of plant species, amounting up to 100 with at least 15 representatives in *Arabidopsis* alone (for a compilation, see Renaudin et al., 1996). Cyclins share a highly homologous region of about 100 amino acids termed the "cyclin box" which is required for their interaction with the CDK catalytic subunit. Analysis of the deduced peptide sequences in the conserved cyclin box has enabled the classification of these cyclins into nine groups: A1, A2, A3, B1, B2, D1, D2, D3, and D4 reflecting their homologies to the mammalian cyclins A, B, and D (Renaudin et al., 1996; Burssens et al., 1998; De Veylder et al., 1999 and references cited therein). See also Table 1 for a list of cyclins relevant for the invention. Cyclins with homology to mammalian Cyclin C have been recently identified in rice and *Arabidopsis*, thus adding even more complexity to the family of cyclins in plants.

The plant A- and B- type cyclins are the so called 'mitotic cyclins' with an important function during mitosis, while the D-type cyclins (so called G1 cyclins) are thought to play a key role at the entry of S phase. The transcriptional regulation of the mitotic cyclins CYCA2;1 and CYCB1;1 of A. thaliana has been analysed in detail in synchronised tobacco BY2-cells. Promoter activity of CYCA2; 1 is switched on upon entry of S phase and persists during G2 phase to be maximal at the end of G2 phase. CYCB1;1 is expressed in a more narrow window of the cell cycle, namely upon exit of S phase and G2 phase with maximal mRNA levels at the entry of mitosis (Shaul et al 1996). Moreover, by developmental expression analysis, the presence of CYCB1;1 transcripts was exclusively linked with actively dividing cells (Ferreira et al., 1994), implying that CYCB1;1 is involved in the regulation of mitosis. Plant D cyclins, by analogy with their animal homologues, have been proposed to control the G1 progression in response to growth factors and nutrients (Dahl et al., 1995; Soni et al., 1995). Moreover ectopic expression has been shown to render cell division in Arabidopsis independent of the growth

4

hormone cytokinin (Riou-Khamlichi et al 1999). Cyclins CYCD2;1 and CYCD3;1 from tobacco are found to be expressed predominantly in G-M (Sorrell et al., 1999), suggesting that D-type cyclins in plants may also be involved in mitotic events. A novel cyclin CYCD4 from Arabidopsis has also been identified and is shown to be expressed during vascular tissue development, embryogenesis, and formation of lateral root primordia (De Veylder et al. 1999).

CDK/Cyclin complexes

There is evidence showing that CDKs and cyclins interact to form functional protein complexes. Bögre et al. (1997) have found that protein fractions from alfalfa extracts corresponding to monomeric CDKs are essentially devoid of kinase activity as measured by histone H1 phosphorylation and, on the other hand, alfalfa protein complexes immunoprecipitated with antibodies against the human cyclin A or alfalfa cyclin CYCB2;2 exhibit appropriate histone H1 kinase activity (Magyar et al., 1993; 1997). Immunolocalization of CDC2Zm and mitotic cyclins in maize suggest the occurrence of several possible combinations of CDKs and cyclins (Mews et al 1997). Further, the inventors have shown, using the two-hybrid system, the interaction of CYCD1;1 (De Veylder et al., 1997a) and CYCD4;1 (De Veylder et al., 1999) with CDC2aAt. Additionally, the inventors are able to purify active kinase complexes from Arabidopsis cells that contain selectively either CDC2aAt or CDC2bAt. The following protein complexes have been purified: CDC2a with CYCB2;2, CDC2a with CYCA2;2, CDC2b with CYCB1;1, CDC2b with CYCA2;2.

Other cell cycle complexes

A number of other cell cycle proteins have been shown to interact with one another to form active complexes. Some complexes relevant to the invention are involved in the initiation of DNA replication and cell division including facilitating the entry into S phase of quiessant cells (see Leatherwood 1998, Helin 1998 for reviews). The complexes include ORC1/CDC6 or CDC7/DBF4 or E2F/DP.

Importance of the cell cycle

The presence of multiple cyclins and CDKs enables the fine regulation of cell cycle controls and checkpoints since different kinase activities are involved at different points within the cell cycle (Burssens et al, 1998 and references cited therein). The importance of the cell cycle for plant growth and development is illustrated by the observed growth inhibition in response to chemical and radiation treatments, that specifically block cell cycle progression (Foard and Haber, 1961b; Ivanov, 1994). Moreover, as in yeast and animal systems (for an overview see e.g. (Murray and Hunt, 1993)), it is expected that the majority of mutations of cell cycle genes are either lethal or result in severe growth reductions. Inversely, if the cell cycle plays a role in plant growth regulation, it is possible to modify plant growth rates by manipulating the expression of cell cycle genes that are limiting cell division and thereby plant growth rates. Indeed, Doerner, 1996 has suggested that ectopic expression of CYCB1;1 under the control of the CDC2a promoter in A. thaliana plants accelerates root growth without altering the pattern of lateral root development or inducing neoplasia. In contrast to this data, the inventors have shown that constitutive overexpression of CYCB1;1 alone does not lead to any significant growth rate differences in at least two independently transformed lines.

CDK activity depends on the interaction with regulatory cyclins and is limiting for cell cycle progression. Different CDK/cyclin complexes act at different time-points of the cell cycle, although different CDK/cyclin complexes may also act at the same time-points of the cell cycle. Complexes of A-type CDKs (such as CDC2a) with D-type cyclins are acting at the G1 phase and are involved in recruiting G0 cells in the G1 phase of the cell cycle. Complexes between A-type CDKs and A-type cyclins are operational in S- and G2 phase. At G2- and M-phase complexes between A-and B-type CDKs and B-type cyclins are controlling the progression of the cell cycle.

6

Two major checkpoints are operational during the cell cycle, one at the G1/S boundary and one at the G2/M boundary. At these checkpoints the activity of the appropriate CDK/cyclin complexes is controlled, either by interaction with inhibitory proteins (at the G1/S transition), or by inhibitory phosphorylations mediated by a Wee1 kinase (at the G2/M transition). Only when the conditions are favourable is the CDK/cyclin kinase activity restored either by inactivation of the inhibitor, or by dephosphorylation of the CDK/cyclin complex through the action of CDC25.

The current invention describes methods to overcome the inhibition of CDK/cyclin activity. We disclose that the simultaneous (over)expression of a CDK and its regulatory cyclin overrides the potential inhibition of the CDK/cyclin complex at the checkpoints. Depending on the specific CDK/cyclin combination different effects are expected in plant growth, yield or architecture.

Thus, the technical problem underlying the present invention is to provide means and methods for enhancement of plant growth, and/or yield and/or modified architecture in particular in the entire plant, or specific parts of said plant, which are particularly useful in agriculture.

The solution to the technical problem is achieved by providing the embodiments characterised in the claims.

SUMMARY OF THE INVENTION

The present invention relates to a method for modifying plant growth and/or yield and/or architecture, in particular modifications to plant growth and development mediated by cell cycle protein complexes, thereby improving the agricultural and commercial value of these plants. Surprisingly, it has been found that the overexpression of at least two proteins forming subunits of a

protein complex in particular cells, tissues or organs of the plant would produce enhanced plant growth and/or yield and/or architecture compared to otherwise non-transformed plants.

7

Accordingly, the present invention relates to a method for modifying plant growth and/or yield and/or modifying architecture comprising introducing into a plant cell, plant tissue or plant (a) nucleic acid molecule(s) or regulatory sequence(s), wherein the introduction of the nucleic acid molecule(s) or regulatory sequence(s) result(s) in an increased or de novo expression of at least two cell cycle interacting proteins capable of forming a (heteromeric) complex in a plant cell.

Surprisingly, it has been found that the simultaneous ectopic expression and/or overexpression of more than one cell cycle interacting protein, of which, preferably, at least one is a protein kinase and at least one is a protein forming a complex with such protein kinase and regulating the activity of said protein kinase, leads to enhanced plant growth and/or yield and/or modified architecture compared to wild type plants.

This effect is surprising in the light of the fact that (i) CDC2a is known to be constitutively expressed throughout the cell cycle and therefore not obviously a rate limiting factor for cell division and (ii) Doerner et al. 1996 have shown growth stimulation based on ectopic overexpression of CYCB1;1 alone. The inventors have shown that, under their experimental conditions, no growth stimulating effect of CYCB1;1 was observed whereas a growth stimulation was observed with simultaneous (over)expression of both CYCB1;1 and CDC2a. Even when one assumes that CYCB1;1 overexpression may have a growth stimulating effect in itself under certain conditions, the current invention demonstrates that it will be advantageous to use co-expression of both CYCB1;1 and CDC2a since this leads to growth stimulation under conditions in which CYCB1;1 alone does not stimulate growth. The combined

8

overexpression may therefore stimulate growth under a wider range of conditions, including environmental conditions, such as high or low availability of water and nutrients, high or low temperature, high or low light, etc.

In a preferred embodiment of this invention, the protein kinase is a cyclin dependent kinase (CDK). In a more preferred embodiment, the CDK is a PSTAIRE type cyclin dependent kinase. In a most preferred embodiment the CDK is a CDC2a. In an alternatively preferred embodiment, the CDK is a B-type cyclin dependent kinase namely CDC2b.

In a further preferred embodiment of this invention the second protein is a cyclin. In a more preferred embodiment the cyclin is a G1 cyclin such as a D-type cyclin (e.g. CYCD4;1) or an E-type cyclin. In an alternatively preferred embodiment of the invention the cyclin is a mitotic cyclin such as a B-type cyclin like CYCB1;1.

In a further preferred embodiment the cell cycle interacting proteins to be coexpressed and forming the relevant cell cycle complex are ORC1 with CDC6 or CDC7 with DBF4 or E2F with DP.

In a further preferred embodiment of the invention both proteins of the cell cycle complex are ectopically expressed under the control of a constitutive promoter such as the 35S promoter. It will be clear to the man skilled in the art that both proteins may also be expressed under the control of other promoters such as tissue specific promoters, which may be the same for both proteins or which may be different as long as those promoters are driving simultaneous expression of both proteins in at least one tissue. Growth stimulation occurs in particular in those tissues in which both proteins are simultaneously (over)expressed.

The man skilled in the art will see various ways of implementing said method in plants.

9

In one embodiment of the invention plants are transformed separately with one protein of the cell cycle complex, e.g. or protein kinase such as CDC2a, and with the other interacting cell cycle protein of the complex such as a regulatory protein of such protein or protein kinase, e.g. CYCD4;1. Such plants are subsequently crossed and the offspring plants that contain both transgenes are selected and demonstrated to exhibit modified growth, yield and/or architectural characteristics in comparison with a wild type plant.

In another embodiment of the invention plants exhibiting simultaneous ectopic expression and/or overexpression of the two genes according to the present invention will be obtained via so called cotransformation. Each gene will be present in a different vector or gene construct (e.g. an *Agrobacterium* vector) and during the transformation step both vectors will be used in combination. The success rate of cotransformation will be highest when both vectors contain a different marker gene (e.g. bar, nptll, hyg,...) and when the selection will be performed with both selective agents; it is also possible to use only one selectable marker gene and its corresponding selective agent and then to identify cotransformants by means of genetic analysis (e.g. PCR based methods). Alternatively each gene will be present on the same vector or gene construct and the plant transformed with such a vector.

In yet another embodiment of the invention specific vectors will be constructed in accordance with the invention. Such vectors will contain a nucleic acid molecule, e.g. a gene, encoding one protein of the cell cycle complex, e.g. encoding a protein kinase, under the control of a given promoter sequence as well as a nucleic acid molecule, e.g. a gene, encoding the other interacting cell cycle protein of the said complex, e.g. regulating the protein kinase activity, under the control of a given promoter sequence. In

addition to the promoters other control sequences may be present. The promoters of both genes may be identical or may be different as long as there is simultaneous expression in at least one tissue. Bidirectional promoters such as the TR promoter may also be used to drive expression of both genes.

The transgenic plants, plant tissues, plant organs or plant cells obtained by the method according to the invention are obtainable from a monocotyledonous plant or dicotyledonous plant.

The invention also relates to a transgenic plant cell comprising an overexpressed protein complex obtainable according to any of the methods of the present invention. A transgenic plant or plant tissue comprising said plant cells and harvestable parts or propagation material of those plants are part of the invention too.

The invention also relates to the vectors necessary to obtain transformed plants in accordance with the previous embodiments of the invention, those vectors are characterized by the fact that they contain both a protein kinase gene and a gene encoding a regulatory protein regulating the activity of said protein kinase.

The invention is also related to utilisation in hybrid seeds in the following way. Two transgenes of interest, each present in a homozygous condition in one of the parents of a hybrid, will be present in combination and in a heterozygous condition in the hybrid seed, thus providing the hybrid seed with the benefit of accelerated growth based on the simultaneous ectopic expression and/or overexpression of the two transgenes. Seed harvested from the F1 hybrid plants will segregate for both transgenes and only 9 out of 16 plants of the F2 generation will possess the two transgenes, thus resulting in additional protection of the value of the hybrid seed.

11

In a still further embodiment the present invention relates to composition comprising the above-described nucleic acid molecules, regulatory sequences or vectors, containing the same identified by the method of the present invention.

In another embodiment the invention relates to the use of the transformed cells or the above described nucleic acid molecules, regulatory sequences or vectors for the production of more biomass, secondary metabolites or additives for plant culturing in plant cell culture.

Detailed description of the invention

The present invention relates to a method for modifying plant growth and/or yield and/or modifying architecture comprising introducing into a plant cell, plant tissue or plant (a) nucleic acid molecule(s) or regulatory sequence(s), wherein the introduction of the nucleic acid molecule(s) or regulatory sequence(s) result(s) in an increased or de novo expression of at least two cell cycle interacting proteins capable of forming a (heteromeric) complex in a plant cell.

The term "growth" is a concept well known to the person skilled in the art and includes increased crop growth and/or enhanced biomass.

The term "modifying plant growth and/or yield" refers to a general alteration in the growth of the plant, its tissues or organs or the yield as examplified below. Preferably, "modifying plant growth" relates to an acceleration, enhancement or promotion of plant growth.

"Architecture" refers to the general morphology of a plant including any one of more structural features including the shape, size, number, colour, texture, arrangement and patternation of any cell, tissue or organ or groups of cells,

tissues, or organs of plants including the root, leaf, shoot, fruit, petiole, trichome, flower, sepals, petal, hypocotyl, stigma, style, stamen, pollen, ovule, seed, embryo, endosperm, seed coat, aleurone, fibre nodule, cambium, wood, heartwood, parenchyma, erenchyma, selve element, phloem, or vascular tissue amongst others.

"Modifying yield" refers to an altered, preferably increased or enhanced biomass of either the total plant or specific tissues or organs of plants such as root, leaf, shoot, fruit, petiole, trichome, flower, sepals, petal, hypocotyl, stigma, style, stamen, pollen, ovule, seed, bulb, embryo, endosperm, seed coat, aleurone, fibre, nodule, cambium, wood, heartwood, parenchyma, erenchyma, seive element, phloem, or vascular tissue. "Yield" also refers to accumulation of metabolites and/or the sink/source relationships in the total plant or specific cells, tissues or organs of the plant such as root, leaf, shoot, fruit, petiole, trichome, flower, sepals, petal, hypocotyl, stigma, style, stamen, pollen, ovule, seed, embryo, endosperm, seed coat, aleurone, fibre, nodule, cambium, wood, heartwood, parenchyma, erenchyma, seive element, phloem, or vascular tissue. This means, for instance that increased growth and/or yield results from increased growth rate or increased root size or shoot growth or, alternatively, in an increased yield because of enhanced fruit growth.

As will be known to those skilled in the art, plants may modify one or more plant growth and/or architectural and/or yield characteristics in response to external stimuli, such as, for example, a plant pathogenic infection, or an external stress or environmental stress (e.g. anoxia, hypoxia, high temperature, low temperatures, light, daylength, drought, flooding, salt stress, dehydration, heavy metal contamination, nutrient/mineral deficiency, amongst others). Accordingly, for the present purpose, it shall be understood that a plant growth or architectural or yield characteristic that has been modified in response to one or more external stimuli is within the scope of the inventive

method described herein, notwithstanding that the imposition of said external stimuli is not an essential feature of the present invention.

The term "simultaneous" as used in the context of "simultaneous ectopic expression and/or overexpression" or "simultaneous (over)expression" shall mean preferably at the same time and in the same cells but at least that, although not expressed at the same time, there is an overlap in the timing of the existence of the proteins of interest so that the proteins are capable of forming a complex.

The term "cell cycle" as used herein shall be taken to include the cyclic biochemical and structural events associated with growth and with division of cells, and in particular with the regulation of the replication of DNA and mitosis. Cell cycle includes phases called: G0 (gap 0), G1 (gap 1), DNA replication (S), G2 (gap 2), and mitosis including cytokinesis (M). Normally these four phases occur sequentially. However, the cell cycle also includes modified cycles such as endomitosis, acytokinesis, polyploidy, polyteny, endopolyploidisation and endoreduplication or endoreplication.

The term "cell cycle interacting protein", "cell cycle protein", or "cell cycle control protein" as denoted herein means a protein which exerts control on or regulates or is required for the cell cycle or part thereof, of a cell, tissue, organ or whole organism and/or DNA replication. It may also be capable of binding to, regulating or being regulated by cyclin dependent kinases or their subunits. The term also includes peptides, polypeptides, fragments, variants, homologues, alleles or precursors (e.g. preproproteins or preproteins) thereof. The cell cycle interacting protein is preferably of plant origin, although it may also be the yeast homologues thereof.

The cell cycle interacting protein is preferably a protein kinase, in particular a cyclin dependent kinase (CDK) for example an A-type (CDC2a) or a B-type (CDC2b) CDK.

The cell cycle interacting protein is also preferably a cyclin, including cyclins A, B, C, D and E, and in particular CYCA1;1, CYCA2;1, CYCA3;1, CYCB1;1, CYCB1;2, CYCB2;2, CYCD1;1, CYCD2;1. CYCD3;1, and CYCD4;1 (Evans et al. 1983; Francis et al. 1998; Labbe et al. 1989; Murray and Kirschner 1989; Renaudin et al 1996; Soni et al 1995; Sorrell et al 1999; Swenson et al 1986). Cyclins are also referred to in Table 1.

The cell cycle interacting protein preferably includes proteins involved in the control of entry and progression through S phase in particular ORC1, CDC6, (Nevins, 1992; Liang, 1995) CDC7, DBF4 kinase, E2F (WO 99/58681; WO 99/53075) and DP (WO 99/53075).

The term "capable of forming a (heteromeric) complex" as used herein means that said at least two cell cycle interacting proteins bind, interact or associate with each other to form a complex in a cellular, preferably plant cellular environment. Preferably, the complex formation in the cell leads to the induction of potential processes of cell division, preferably cell proliferation. Examples of such cell cycle interacting proteins include, but are not limited to protein kinases e.g., cyclin-dependent kinases (CDKs), and their activating associated subunits, namely cyclins (CYCs). Other cell cycle interacting proteins capable of forming a (heteromeric) complex include ORC1 and CDC6, CDC7 and DBF4, and E2F and DP.

The resulting complex of at least two cell cycle interacting proteins capable of forming a (heteromeric) complex is termed a "cell cycle complex". A preferred cell cycle complex is a CDK/cyclin complex, namely the complex formed when a preferably functional cyclin associates with its appropriate CDK,

15

preferably a functional form thereof. Such complexes may be active in phosphorylating proteins and may or may not contain additional protein species. Alternatively preferred cell cycle complexes are ORC1/CDC6 or CDC7/DBF4 or E2F/DP.

The invention includes modified forms of a cell cycle interacting protein, including homologues or analogues (as defined below) thereof. A preferred modified cell cycle interacting protein is a modified CDK protein wherein the modification removes the inhibitory effect of phosphates on CDC2, in particular the threonine-14 and/or tyrosine-15 have been substituted with non-phosphorylatable residues such as phenylalanine and/or alanine. Examples of such modified CDKs include Cdc2aA14F15 and Cdc2bA14F15. Another form of modification includes the mutation of the amino acid residue responsible for ATP binding, namely the D residue is replaced with an N residue to form for example Cdc2b.N161, Cdc2f.N164, or Cdc2aN147.

Another preferred modified cell cycle interacting protein is a modified cyclin protein wherein the modification results in the stablisation of the cyclin. Such modification may be the result of the mutation or complete or partial removal of the destruction box (D box) motif (RxxLxx[L/I]xN) in the cyclin N-terminal domain (where R and L residues are highly conserved and x stands for any amino acid) (Plesse et al 1998). Examples of modified cyclins for use in the present invention include CYCA2;2 Δ 64, CYCA2;3 Δ 63, CYCB2;1 Δ 44 (where Δ – is the truncated form lacking the N-terminal part containing the destruction box).

"Homologues" of cell cycle interacting proteins are those peptides, oligopeptides, polypeptides, proteins and enzymes which contain amino acid substitutions, deletions and/or additions relative to a non-mutant or wild-type cell cycle interacting protein polypeptide, without altering one or more of its cell cycle control properties. To produce such homologues of cell cycle

16

interacting protein, amino acids present in the protein can be replaced by other amino acids having similar properties, for example hydrophobicity, hydrophobic moment, antigenicity, propensity to form or break helical structures or sheet structures, and so on.

Substitutional variants are those in which at least one residue in the cell cycle interacting protein amino acid sequence has been removed and a different residue inserted in its place. Amino acid substitutions are typically of single residues, but may be clustered depending upon functional constraints placed upon the polypeptide; insertions will usually be of the order of about 1-10 amino acid residues and deletions will range from about 1-20 residues. Preferably, amino acid substitutions will comprise conservative amino acid substitutions, such as those described *supra*.

Insertional amino acid sequence variants are those in which one or more amino acid residues are introduced into a predetermined site in the cell cycle interacting protein. Insertions can comprise amino- terminal and/or carboxyl terminal fusions as well as intra-sequence insertions of single or multiple amino acids. Generally, insertions within the amino acid sequence will be smaller than amino or carboxyl terminal fusions, of the order of about 1 to 4 residues.

Deletional variants are characterised by the removal of one or more amino acids from the cell cycle interacting protein sequence.

Amino acid variants of the cell cycle interacting polypeptide may readily be made using peptide synthetic techniques well known in the art, such as solid phase peptide synthesis and the like, or by recombinant DNA manipulations. The manipulation of DNA sequences to produce variant proteins which manifest as substitutional, insertional or deletional variants are well known in the art. For example, techniques for making substitution mutations at

predetermined sites in DNA having known sequence are well known to those skilled in the art, such as by M13 mutagenesis or other site-directed mutagenesis protocol.

"Analogues" of a cell cycle interacting protein are defined as those peptides, oligopeptides, polypeptides, proteins and enzymes which are functionally equivalent to a cell cycle interacting protein.

Analogues of a cell cycle interacting protein include those CDKs, cyclins etc and modified versions thereof that comprise peptides, polypeptides, proteins, and enzymes that are capable of functioning in a plant cell and/or plant tissue and/or plant organ and/or whole plant to produce the same modified plant growth and/or yield and/or architectural characteristics as the ectopic expression and/or (over)expression of such cell cycle interacting proteins forming a cell cycle complex.

The terms "gene(s)", "polynucleotide", "nucleic acid sequence", "nucleotide sequence", "DNA sequence" or "nucleic acid molecule(s)" as used herein refers to a polymeric form of nucleotides of any length, either ribonucleotides or deoxyribonucleotides. This term refers only to the primary structure of the molecule. Thus, this term includes double- and single-stranded DNA, and RNA. It also includes known types of modifications, for example, methylation, "caps" substitution of one or more of the naturally occurring nucleotides with an analog.

The present invention denotes nucleic acid molecules which enhances expression of said cell cycle interacting proteins. For example, said nucleic acid molecules comprise a coding sequence of a mentioned protein or of a regulatory protein, e.g., a transcription factor, capable of inducing the expression of said cell cycle interacting protein.

The term "regulatory sequence" as used herein denotes a nucleic acid molecule increasing the expression of the said protein(s), e.g. of cell cycle interacting protein(s), due to its integration into the genome of a plant cell in close proximity to the gene(s), e.g. encoding said cell cycle interacting protein(s). Such regulatory sequences comprise promoters, enhancers, inactivated silencer intron sequences, 3'UTR and/or 5'UTR coding regions, protein and/or RNA stabilizing elements or other gene expression control elements which are known to activate gene expression and/or increase the amount of the gene products.

The introduction of said nucleic acid molecule(s) leads to de novo expression, or if the mentioned regulatory sequence(s) is used to increase in expression of said proteins, resulting in an increased amount of active protein in the cell. Thus, the present invention is aiming at providing de novo and/or increased activity of e.g., cell cycle interacting proteins.

The experiments performed underlying the current invention clearly show that overexpression of *CYCB1;1* and *CDC2a* in conjunction results in a growth stimulation of both root and shoot of between 10% to 30%. This growth stimulation requires the overexpression of both genes, since there was no growth stimulating effect of overexpression of either gene alone; and in the case of *CDC2a* overexpression alone there is a growth reduction. The relative reduction in growth rate of the roots in the CDC2a overexpressing lines increases in function of time. Growth rates 3 days after sowing were similar.

F1 seedlings overexpressing both CYCB1;1 and CDC2a exhibited increased growth. This increase was apparent 3 days after sowing (earliest measurement) throughout the entire observation period.

It was found that the growth increase is not due to more rapid germination of the seeds, since no difference in timing of germination was observed and the general growth rate pattern with a flattening of the curve at 11 days after germination was maintained.

19

Further, it was found that increased cell number rather than increased cell size explains the observed increase in growth (see figure 6).

Furthermore, support has been found that the increased cell number is due to a more rapid cell division rate rather than to a larger number of dividing cells in the meristem. This explanation is coherent with a role of the protein kinase and its regulatory protein in the control of cell division.

Whereas in the current invention, transgenic lines overexpressing both CDC2a and CYCB1;1 were obtained by crossing a line homozygous for a 35ScaMV-CDC2a construct with a line homozygous for a modified 35S-CYCB1;1 construct, it is clear for a person skilled in the art that the same effect could also be obtained by introducing in one plant e.g., a DNA construct in which both the CDC2a and CYCB1;1 are placed under a constitutive or tissue specific promoter.

As is evident from the above, in one embodiment of the method of the present invention said nucleic acid molecule(s) encode(s) said cell cycle interacting protein(s) and the regulatory sequence(s) is (are) capable of increasing the expression of a gene encoding said cell cycle interacting protein(s). This means, that a nucleic acid molecule comprises a coding sequence for a cell cycle interacting protein as defined herein.

A "coding sequence" is a nucleotide sequence which is transcribed into mRNA and/or translated into a polypeptide when placed under the control of appropriate regulatory sequences. The boundaries of the coding sequence are determined by a translation start codon at the 5'-terminus and a translation stop codon at the 3'-terminus. A coding sequence can include, but is not limited to mRNA, cDNA, recombinant nucleotide sequences or genomic DNA, while introns may be present as well under certain circumstances.

As has been demonstrated in the appended examples, one of said cell cyclin interacting proteins is a protein kinase. Therefore, in a preferred embodiment

20

of the method of the present invention one of said cell cycle interacting proteins is a protein kinase.

In a particularly preferred embodiment of the method of the present invention said protein kinase is a cyclin-dependent kinase (CDK) and the other of said cell cycle interacting proteins is a cyclin. CDKs and cyclins that can be employed according to the invention are described in Table 1, in Segers et al., 1997 or Renaudin et al., 1996, (the disclosure contents of which are hereby incorporated by reference). Preferred coexpression of combinations of CDKs and cyclins in a cell cycle complex and the resulting phenotype are described below:

- 1. A-type CDK with a cyclin D2 or cyclin D4: Without being bound by any theory or mode of action overexpression of A-type CDKs and cyclin D2 and/or cyclin D4 shortens the G1 phase and overrides the checkpoints, which monitors the availability of sugars. As such plants complete their lifecycle (from seed to seed) faster. It also is expected that plants can complete extra rounds of cell division resulting in an enhanced production of biomass. By using meristem specific promoters (e.g. promoters active in root- or shoot meristems, or subdomains thereof, promoters active during early seed development, cambium specific promoters etc.) growth of particular organs can be enhanced.
- 2. A-type CDK with a cyclin D4: Without being bound by any theory or mode of action overexpression of A-type CDKs and cyclin D4 elevates the threshold that cells needs to re-enter the cell cycle. As such plants cells are more easily regenerated and form more structures (such as lateral roots) of which the formation is dependent on the re-activation of the cell cycle (G0 to G1 transition).

- 3. A-type CDK with a D-type cyclin: The G1/S checkpoint as well as the G2/M checkpoint is of importance for the arrest of cell division under environmental stress conditions. It is anticipated that overexpression of A-type CDKs and D-type cyclins would result in plants with growth characteristics that are more tolerant to stress conditions which cause a cell cycle arrest at the G1/S boundary.
- 4. A-type CDK with a B-type cyclin or a B-type CDK with a B-type cyclin: overexpression of A-type and/or B-type CDKs with B-type cyclins overrides the G2/M checkpoints. Such plants are expected to have growth characteristics less sensitive to environmental stress conditions, such as osmotic stress and complete faster the G2 phase. Furthermore, cells will become less dependent upon the continuous availability of cytokinins.
- 5. A-type CDK with a D-type cyclin: Without being bound by any theory or mode of action the G1 phase is thought to be of importance for growth control of plant cells and an arrest of cells at the G1 phase (e.g. through the use of a dominant negative A-type CDK) cause cells to become larger. As such it is anticipated that co-expression of A-type CDKs and D-type cyclins enhances the progression through the G1 phase and reduces the average cell size. More cells per unit surface results in a modification of the tissue texture (relative more cell wall material, more membranes etc.) and has an important impact on quality traits. For example the difference between spring and summer wood in trees is largely a consequence of differences in cell size. Possibly fibrelength of cells (e.g. cotton fibres) can be modified through an alteration of the length of the G1 phase.
- 6. B-type CDK with an A-type cyclin: plant A-type cyclins are expressed from mid S till early M phase, therefore it can be expected that the co-expression of CDC2bAt with CYCA results in an enhanced progression though these cell cycle phases. Also cells may become less sensitive to the G2/M checkpoint

control, making the plants putatively less sensitive to stress conditions and plant growth regulators which operate at this transition point. The result would be that these plants grow better in suboptimal conditions, compared to normal plants, noting that suboptimal conditions may occur frequently even under good agronomical conditions. A higher yield is anticipated under most, but particularly under suboptimal/marginal, agricultural conditions.

The following Table 2 also illustrates preferred CDK/cyclin cell cycle complexes for use in the performance of the application:

TABLE 2

CYCLIN	CDK
CYCA2;1	Cdc2a
CYCA2;2	Cdc2a, Cdc2b, Cdc2f,
	Cdc2bN161 Cdc2aN146
CYCA2;3	Cdc2b, Cdc2f, Cdc2bN161,
	Cdc2aN146
CYCB1;1	Cdc2a, Cdc2b
CYCB1;2	Cdc2a, Cdc2b
CYCB2;1	Cdc2b; Cdc2f
CYCB2;2	Cdc2a, Cdc2b
CYCD1;1	G1-CDK, Cdc2a
CYCD2;1	G1-CDK, Cdc2a
CYCD3;1	G1-CDK, Cdc2a
CYCD4;1	Cdc2a; Cdc2b, Cdc2f,
	Cdc2bN161 Cdc2aN146,
	Cdc2fN164

It may not be necessary to express or induce the expression of native wild type cell cycle interacting proteins, such as above described CDKs and cyclins but it may be sufficient to provide for complex formation of at least a catalytic and/or regulatory subunit of said cell cycle interacting proteins. Therefore, another embodiment of the method of the present invention said nucleic acid molecule(s) encode(s) at least a catalytic and/or regulatory subunit of said cell cycle interacting protein(s).

PCT/EP00/02441

Components of CDK/cyclin complexes that can be employed in accordance with the method of the present invention and how to obtain them are known to the person in skilled and are described, e.g., in WO 98/41642, WO 92/09685 the disclosure of which is hereby incorporated by reference.

One aspect of the invention provides a method of modifying plant growth and/or yield and/or architecture by expressing in particular cells, tissues or organs of a plant, at least two subunits of a cell cycle complex operably under the control of a regulatable promoter sequence selected from the list comprising cell-specific promoter sequences, tissues-specific promoter sequences and organ-specific promoter sequences. Examples of such promoters include promoters which are:

- stem-expressible and more specifically in the stem cambium: to increase strength and thickness of a plant stem to confer improved stability and wind-resistance on the plant
- meristem expressible: to inhibit or reduce apical dominance or increasing the bushiness of a plant. This is a desirable phenotype in a number of crop plants for example in the different Brassica species.
- tuber expressible: to increase or improve tuber production in the plant
- **seed** expressible: to increase seed production in plants in particular to increase seed set and/or seed production and/or seed yield.
- endosperm expressible: those skilled in the art will be aware that grain
 yield in crop plants is largely a function of the amount of starch produced
 in the endosperm of the seed. The amount of protein produced in the
 endosperm is also a contributing factor to grain yield. In contrast that
 embryo and aleurone layers contribute little in terms of the total weight of
 the mature grain. Therefore endosperm-expressible promoters provide the
 advantage of increasing grain size and grain yield of the plant.
- root expressible: to increase or enhance the production of roots or storage organs derived from roots
- nodule expressible: to increase the nitrogen-fixing capability of a plant.

- embryo expressible: embryo size being important for growth after germination
- leaf expressible
- flower expressible
- fruit expressible

Examples of promoters suitable for use in gene constructs of the present invention include those listed in Table 3, amongst others. The promoters listed in Table 3 are provided for the purposes of exemplification only and the present invention is not to be limited by the list provided therein. Those skilled in the art will readily be in a position to provide additional promoters that are useful in performing the present invention. Table 4 describes constitutive promoters for use in the present invention.

Preferred promoters may contain additional copies of one or more specific regulatory elements, to further enhance expression and/or to alter the spatial expression and/or temporal expression of a nucleic acid molecule to which it is operably connected. For example, copper-responsive, glucocorticoid-responsive or dexamethasone-responsive regulatory elements may be placed adjacent to a heterologous promoter sequence driving expression of a nucleic acid molecule to confer copper inducible, glucocorticoid-inducible, or dexamethasone-inducible expression respectively, on said nucleic acid molecule.

In case the above-described cell cycle interacting proteins or at least one of them are to be expressed de novo, it is preferred to employ in the method of the present invention genes encoding such cell cycle interacting proteins, wherein said gene is expressible in plant cells. Thus, in another embodiment the method of the present invention said nucleic acid molecule(s) is (are) operatively linked to control sequences allowing the expression of the nucleic acid molecule(s) in the plant. Said control sequences comprise a promoter,

enhancer, silencer, intron sequences, 3'UTR and/or 5'UTR regions, protein and/or RNA stabilizing elements. Preferably, said control sequence is a chimeric, tissue specific, constitutive or inducible promoter.

Preferably, both proteins are expressed under the control of a promoter which is active in non differentiated plant cells or in plant protoplasts growing in an artificial medium. The increased growth rate of the cells results in increasing growth of the plant cells in plant cell culture, thus allowing the production of more biomass in plant cell culture. Plant cell production in plant culture can be useful for the production of certain secondary metabolites of plants which may be useful in the pharmaceutical, cosmetics, food industry etc.

The present invention further relates to a nucleic acid molecule encoding at least two cell cycle interacting proteins as mentioned above.

The present invention also relates to vectors, particularly plasmids, cosmids, viruses, bacteriophages and other vectors used conventionally in genetic engineering that comprise the nucleic acid molecule or at least two nucleic acid molecules and/or regulatory sequences according to the invention. Methods which are well known to those skilled in the art can be used to construct various plasmids and vectors; see, for example, the techniques described in Sambrook, Molecular Cloning A Laboratory Manual, Cold Spring Harbor Laboratory (1989) N.Y. and Ausubel, Current Protocols in Molecular Biology, Green Publishing Associates and Wiley Interscience, N.Y. (1989), (1994). Plasmids and vectors to be preferably employed in accordance with the present invention include those well known in the art. Alternatively, the nucleic acid molecules and vectors of the invention can be reconstituted into liposomes for delivery to target cells.

In a preferred embodiment the vector of the present invention comprises separate nucleic acid molecules encoding at least one of said cell cycle interacting proteins.

In a preferred embodiment the nucleic acid molecule present in the vector is linked to (a) control sequence(s) which allow the expression of the nucleic acid molecule or of cell cycle interacting proteins in a host cell, e.g. prokaryotic and/or eukaryotic cells.

The term "control sequence" refers to regulatory DNA sequences which are necessary to effect the expression of coding sequences to which they are ligated. The nature of such control sequences differs depending upon the host organism. In prokaryotes, control sequences generally include promoter, ribosomal binding site, and terminators. In eukaryotes generally control sequences include promoters, terminators and, in some instances, enhancers, transactivators or transcription factors. The term "control sequence" is intended to include, at a minimum, all components the presence of which are necessary for expression, and may also include additional advantageous components. Preferably, said control sequence comprises a constitutive, chimeric, tissue specific or inducible promoter.

The term "operably linked" refers to a juxtaposition wherein the components so described are in a relationship permitting them to function in their intended manner. A control sequence "operably linked" to a coding sequence is ligated in such a way that expression of the coding sequence is achieved under conditions compatible with the control sequences. In case the control sequence is a promoter, it is obvious for a skilled person that double-stranded nucleic acid is used.

Thus, the vector of the invention is preferably an expression vector. An "expression vector" is a construct that can be used to transform a selected host cell and provides for expression of a coding sequence in the selected host. Expression vectors can for instance be cloning vectors, binary vectors or integrating vectors. Expression comprises transcription of the nucleic acid

molecule preferably into a translatable mRNA. Regulatory elements ensuring expression in prokaryotic and/or eukaryotic cells are well known to those skilled in the art. In the case of eukaryotic cells they comprise normally promoters ensuring initiation of transcription and optionally poly-A signals ensuring termination of transcription and stabilization of the transcript, for example, those of the 35S RNA from Cauliflower Mosaic Virus (CaMV). Other promoters commonly used are the polyubiquitin promoter, and the actin promoter for ubiquitous expression. The termination signals usually employed are from the Nopaline Synthase promoter or from the CAMV 35S promoter. A plant translational enhancer often used is the tobaccos mosaic virus (TMV) omega sequences, the inclusion of an intron (Intron-1 from the Shrunken gene of maize, for example) has been shown to increase expression levels by up to 100-fold. (Mait, Transgenic Research 6 (1997), 143-156; Ni, Plant Journal 7 (1995), 661-676). Additional regulatory elements may include transcriptional as well as translational enhancers. Advantageously, the abovedescribed vectors of the invention comprises a selectable and/or scorable marker. Selectable marker genes useful for the selection of transformed plant cells, callus, plant tissue and plants are well known to those skilled in the art and comprise, for example, antimetabolite resistance as the basis of selection for dhfr, which confers resistance to methotrexate (Reiss, Plant Physiol. (Life Sci. Adv.) 13 (1994), 143-149); npt, which confers resistance to the aminoglycosides neomycin, kanamycin and paromycin (Herrera-Estrella, EMBO J. 2 (1983), 987-995) and hygro, which confers resistance to hygromycin (Marsh, Gene 32 (1984), 481-485). Additional selectable genes have been described, namely trpB, which allows cells to utilize indole in place of tryptophan; hisD, which allows cells to utilize histinol in place of histidine (Hartman, Proc. Natl. Acad. Sci. USA 85 (1988), 8047); mannose-6phosphate isomerase which allows cells to utilize mannose (WO 94/20627) and ODC (ornithine decarboxylase) which confers resistance to the ornithine decarboxylase inhibitor, 2-(difluoromethyl)-DL-ornithine, DFMO (McConlogue, 1987, In: Current Communications in Molecular Biology, Cold Spring Harbor

Laboratory ed.) or deaminase from Aspergillus terreus which confers resistance to Blasticidin S (Tamura, Biosci. Biotechnol. Biochem. 59 (1995), 2336-2338).

Useful scorable marker are also known to those skilled in the art and are commercially available. Advantageously, said marker is a gene encoding luciferase (Giacomin, Pl. Sci. 116 (1996), 59-72; Scikantha, J. Bact. 178 (1996), 121), green fluorescent protein (Gerdes, FEBS Lett. 389 (1996), 44-47) or ß-glucuronidase (Jefferson, EMBO J. 6 (1987), 3901-3907). This embodiment is particularly useful for simple and rapid screening of cells, tissues and organisms containing a vector of the invention.

As it is immediately evident to the person skilled in the art, the vectors used according to a method of the present invention can carry nucleic acid molecules encoding the above-mentioned enzymes or enzymatical fragments thereof and fusions of targeting signals to these molecules. The same applies to the above described plant cells, plant tissue and plants transformed therewith. Likewise, said nucleic acid molecules may be under the control of the same regulatory elements or may be separately controlled for expression. In this respect, the person skilled in the art will readily appreciate that the nucleic acid molecules encoding e.g. the domains of cell cycle interacting protein(s) can be expressed in the form of a single mRNA as transcriptional and optionally translational fusions. This means that domains are produced as separate polypeptides or in the latter option as a fusion polypeptide that is further processed into the individual proteins, for example via a cleavage site for proteinases that has been incorporated between the amino acid sequences of both proteins. The resultant protein domains can then selfassemble in vivo. Of course, the domains may also be expressed as a bi- or multifunctional polypeptide, preferably disposed by a peptide linker which advantageously allows for sufficient flexibility of both proteins. Preferably said peptide linker comprises plural, hydrophilic, peptide-bonded amino acids of a length sufficient to span the distance between the C-terminal end of one of said proteins and the N-terminal end of the other of said proteins when said polypeptide assumes a conformation suitable for biological activity of both proteins when disposed in aqueous solution in the plant cell. Examples of the above-described expression strategies can be found in the literature, e.g., for dicistronic mRNA (Reinitiation) in Hefferon (1997), Brinck-Peterson (1996) and Hotze (1995); bifunctional proteins are discussed in Lamp (1998) and Dumas (1997) and for linker peptide and protease refer to Doskeland (1996).

The present invention furthermore relates to host cells comprising a vector as described above or the mentioned complex overexpressed in a plant cell according to the invention wherein the nucleic acid molecule is foreign to the host cell.

By "foreign" it is meant that the nucleic acid molecule is either heterologous with respect to the host cell, this means derived from a cell or organism with a different genomic background, or is homologous with respect to the host cell but located in a different genomic environment than the naturally occurring counterpart of said nucleic acid molecule. This means that, if the nucleic acid molecule is homologous with respect to the host cell, it is not located in its natural location in the genome of said host cell, in particular it is surrounded by different genes. In this case the nucleic acid molecule may be either under the control of its own promoter or under the control of a heterologous promoter. The vector or nucleic acid molecule according to the invention which is present in the host cell may either be integrated into the genome of the host cell or it may be maintained in some form extrachromosomally.

The host cell can be any prokaryotic or eukaryotic cell, such as bacterial, insect, fungal, plant or animal cells. Preferred fungal cells are, for example, those of the genus Saccharomyces, in particular those of the species S. cerevisiae.

In another embodiment of the present invention, a composition comprising vectors wherein each vector contains at least one nucleic acid molecule encoding at least one cell cycle interacting protein is disclosed. The expression of said vectors results in the production of at least two cell cycle interacting proteins and assembly of the same in a complex in vitro or in vivo.

Another object of the invention is a method for the preparation of a cell cycle complex which comprises the cultivation of host cells according to the invention which, due to the presence of a vector or a nucleic acid molecule according to the invention, are able to express such a cell cycle complex, under conditions which allow expression of the cell cycle complex and recovering of the so-produced complex from the culture.

Accordingly, the present invention relates to a cell cycle complex obtainable by said method of the present invention or encodable by the nucleic acid molecule of the present invention.

The term "expression" means the production of a protein or nucleotide sequence in the cell. It includes transcription into an RNA product, post-transcriptional modification and/or translation to a protein product or polypeptide from a DNA encoding that product, as well as possible post-translational modifications. The terms "protein" and "polypeptide" used in this application are interchangeable. "Polypeptide" refers to a polymer of amino acids (amino acid sequence) and does not refer to a specific length of the molecule. Thus peptides and oligopeptides are included within the definition of polypeptide. This term does also refer to or include post-translational modifications of the polypeptide, for example, glycosylations, acetylations, phosphorylations and the like. Included within the definition are, for example, polypeptides containing one or more analogs of an amino acid (including, for example, unnatural amino acids, etc.), polypeptides with substituted linkages,

as well as other modifications known in the art, both naturally occurring and non-naturally occurring.

Methods for the introduction of foreign DNA into plants are also well known in the art. These include, for example, the transformation of plant cells or tissues with T-DNA using Agrobacterium tumefaciens or Agrobacterium rhizogenes, the fusion of protoplasts, direct gene transfer (see, e.g., EP-A 164 575), injection, electroporation, biolistic methods like particle bombardment, pollenmediated transformation, plant RNA virus-mediated transformation, liposomemediated transformation, transformation using wounded or enzyme-degraded immature embryos, or wounded or enzyme-degraded embryogenic callus and other methods known in the art. The vectors used in the method of the invention may contain further functional elements, for example "left border"and "right border"-sequences of the T-DNA of Agrobacterium which allow for stably integration into the plant genome. Furthermore, methods and vectors are known to the person skilled in the art which permit the generation of marker free transgenic plants, i.e. the selectable or scorable marker gene is lost at a certain stage of plant development or plant breeding. This can be achieved by, for example co-transformation (Lyznik, Plant Mol. Biol. 13 (1989), 151-161; Peng, Plant Mol. Biol. 27 (1995), 91-104) and/or by using systems which utilize enzymes capable of promoting homologous recombination in plants (see, e.g., WO97/08331; Bayley, Plant Mol. Biol. 18 (1992), 353-361); Lloyd, Mol. Gen. Genet. 242 (1994), 653-657; Maeser, Mol. Gen. Genet. 230 (1991), 170-176; Onouchi, Nucl. Acids Res. 19 (1991), 6373-6378). Methods for the preparation of appropriate vectors are described by, e.g., Sambrook (Molecular Cloning; A Laboratory Manual, 2nd Edition (1989), Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY). Suitable strains of Agrobacterium tumefaciens and vectors as well as transformation of Agrobacteria and appropriate growth and selection media are well known to those skilled in the art and are described in the prior art (GV3101 (pMK90RK), Koncz, Mol. Gen. Genet. 204 (1986), 383-396; C58C1

(pGV 3850kan), Deblaere, Nucl. Acid Res. 13 (1985), 4777; Bevan, Nucleic. Acid Res. 12(1984), 8711; Koncz, Proc. Natl. Acad. Sci. USA 86 (1989), 8467-8471; Koncz, Plant Mol. Biol. 20 (1992), 963-976; Koncz, Specialized vectors for gene tagging and expression studies. In: Plant Molecular Biology Manual Vol 2, Gelvin and Schilperoort (Eds.), Dordrecht, The Netherlands: Kluwer Academic Publ. (1994), 1-22; EP-A-120 516; Hoekema: The Binary Plant Vector System, Offsetdrukkerij Kanters B.V., Alblasserdam (1985), Chapter V, Fraley, Crit. Rev. Plant. Sci., 4, 1-46; An, EMBO J. 4 (1985), 277-287). Although the use of Agrobacterium tumefaciens is preferred in the method of the invention, other Agrobacterium strains, such as Agrobacterium rhizogenes, may be used, for example if a phenotype conferred by said strain is desired. Methods for the transformation using biolistic methods are well known to the person skilled in the art; see, e.g., Wan, Plant Physiol. 104 (1994), 37-48; Vasil, Bio/Technology 11 (1993), 1553-1558 and Christou (1996) Trends in Plant Science 1, 423-431. Microinjection can be performed as described in Potrykus and Spangenberg (eds.), Gene Transfer To Plants. Springer Verlag, Berlin, NY (1995).

The transformation of most dicotyledonous plants is possible with the methods described above. But also for the transformation of monocotyledonous plants several successful transformation techniques have been developed. These include the transformation using biolistic methods as, e.g., described above as well as protoplast transformation, electroporation of partially permeabilized cells, introduction of DNA using glass fibers, Agrobacterium mediated transformation etc.

Plants may also be transformed by an *in planta* method using *Agrobacterium tumefaciens* such as that described by Bechtold *et al.*, (1993) or Clough *et al* (1998), wherein *A. tumefaciens* is applied to the outside of the developing flower bud and the binary vector DNA is then introduced to the developing microspore and/or macrospore and/or the developing seed, so as to produce a transformed seed without the exogenous application of cytokinin and/or

33

gibberellin. Those skilled in the art will be aware that the selection of tissue for use in such a procedure may vary, however it is preferable generally to use plant material at the zygote formation stage for *in planta* transformation procedures.

The term "transformation" as used herein, refers to the transfer of an exogenous polynucleotide into a host cell, irrespective of the method used for the transfer. The polynucleotide may be transiently or stably introduced into the host cell and may be maintained non-integrated, for example, as a plasmid, or alternatively, may be integrated into the host genome. The resulting transformed plant cell or plant tissue can then be used to regenerate a transformed plant in a manner known by a skilled person.

In general, the plants which can be modified according to the invention and which either show overexpression of a cell cycle complex according to the invention can be derived from any desired plant species. They can be monocotyledonous plants or dicotyledonous plants, preferably they belong to plant species of interest in agriculture, wood culture or horticulture interest, such as a crop plant, root plant, oil producing plant, wood producing plant, agricultured bioticultured plant, fodder or forage legume, companion plant, or horticultured plant e.g., such a plant is wheat, barley, maize, rice, carrot, sugar beet, cichorei, cotton, sunflower, tomato, cassava, grapes, soybean, sugar cane, flax, oilseed rape, tea, canola, onion, asparagus, carrot, celery, cabbage, lentil, broccoli, cauliflower, brussel sprout, artichoke, okra, squash, kale, collard greens or potato.

Thus, the present invention also relates to transgenic plants and plant tissue comprising transgenic plant cells according to the invention. Due to the (over)expression of the protein complex of the invention, e.g., at developmental stages and/or in plant tissue in which they do not naturally occur or are present at low levels, these transgenic plants may show various growth, yield or

architectural modifications in comparison to wild-type plants. In other words, in one embodiment the present invention relates to a transgenic plant cell displaying de novo expressed cell cycle interacting protein complex or an increased amount of said complex compared to a corresponding wild type plant cell. Said transgenic plant cell comprises at least one nucleic acid molecule or regulatory sequence as defined above or obtainable by the method of the present invention. Furthermore, the present invention relates to transgenic plants and plant tissue obtainable by the method of the present invention. As mentioned above, said transgenic plants may display various idiotypic modifications, preferably display modified and/or accelerated and/or enhanced plant growth, root growth and/or yield compared to the corresponding wild type plant.

Preferred characteristics of the transgenic plants in the present invention is for example that the displays and increased cell division rate. In view of the general teaching of the present invention, it will be appreciated that the present invention contemplates the broad application of the inventive method to the modification of a range of cellular processes, including but not limited to the initiation, promotion, stimulation or enhancement of cell division, seed development, tuber formation, shoot initiation, leaf initiation, root growth, the inhibition of apical dominance etc.

Any transformed plant obtained according to the invention can be used in a conventional breeding scheme or in *in vitro* plant propagation to produce more transformed plants with the same characteristics and/or can be used to introduce the same characteristic in other varieties of the same or related species. Such plants are also part of the invention. Seeds obtained from the transformed plants genetically also contain the same characteristic and are part of the invention. As mentioned before, the present invention is in principle applicable to any plant and crop that can be transformed with any of the transformation method known to those skilled in the art and includes for

instance corn, wheat, barley, rice, oilseed crops, cotton, tree species, sugar beet, cassava, tomato, potato, numerous other vegetables or fruits.

In yet another aspect, the invention also relates to harvestable parts and to propagation material of the transgenic plants according to the invention which either contain transgenic plant cells over-expressing the protein complex according to the invention. Harvestable parts can be in principle any useful parts of a plant, for example, flowers, pollen, seedlings, tubers, leaves, stems, fruit, seeds, roots etc. Propagation material includes, for example, seeds, fruits, cuttings, seedlings, tubers, rootstocks etc.

Furthermore, the present invention relates to the use of the above described nucleic acid molecules, regulatory sequences, and vectors for increasing cell division rates in plants, plant cells or plant tissue. Preferably, said increased cell division rates result in increased biomass, plant growth, root and/or shoot growth, increased seed setting. Preferably, said increased cell division rates result in increased plant growth, modified architecture and/or yield, e.g. of harvestable material, which is displayed for instance by (but not limited to) increased or enhanced biomass, root growth, shoot growth, seed set, seed production, grain yield, fruit size, nitrogen fixing capacity, nodule size, tuber formation, stem thickness, endosperm size, number of fruit per plant etc.

The method of the present invention provides plant cells, plant tissue and plants with novel phenotypes due to the increased or de novo formation of complexes of cell cycle interacting proteins. The plants, plant tissue and plant cells of the present invention will allow the understanding of function of cell cycle protein complexes during this cell division may also open up the way for finding compounds that interfere with formation of such complexes. Thus, the present invention provide a basis for the development of mimetic compounds that may be inhibitors or regulators of cell cycle interacting proteins or their encoding genes. It will be appreciated that the present invention also provides

36

cell based screening methods that allow a high-throughput-screening (HTS) of compounds that may be candidates for such inhibitors and regulators.

In a further embodiment the present invention relates to a composition comprising the nucleic acid molecule, the plant cell or the vector of the present invention or the mentioned vector comprised in the composition of the present invention or the mentioned nucleic acid molecules or regulatory sequences.

Further, in one embodiment the present invention relates to the use of the mentioned nucleic acid molecule or the mentioned regulatory sequence or the nucleic acid molecule, the vector or the plant cell of the present invention or the mentioned vectors of the composition of the present invention for the production of more biomass, of secondary metabolites or additives for plant culturing in plant cell culture.

These and other embodiments are disclosed and encompassed by the description and examples of the present invention. Further literature concerning any one of the methods, uses and compounds to be employed in accordance with the present invention may be retrieved from public libraries, using for example electronic devices. For example the public database "Medline" may be utilized which is available on the Internet, for example under http://www.ncbi.nlm.nih.gov/PubMed/medline.html. Further databases and addresses, such as http://www.ncbi.nlm.nih.gov/, http://www.infobiogen.fr/, http://www.fmi.ch/biology/research_tools.html, http:// www.tigr.org/, are known to the person skilled in the art and can also be obtained using, e.g., http://www.lycos.com. An overview of patent information in biotechnology and a survey of relevant sources of patent information useful for retrospective searching and for current awareness is given in Berks, TIBTECH 12 (1994), 352-364.

The present invention is further illustrated by reference to the following nonlimiting examples.

Unless stated otherwise in the examples, all recombinant DNA techniques are performed according to protocols as described in Sambrook et al. (1989), Molecular Cloning: A Laboratory Manual. Cold Spring Harbor Laboratory Press, NY or in Volumes 1 and 2 of Ausubel et al. (1994), Current Protocols in Molecular Biology, Current Protocols. Standard materials and methods for plant molecular work are described in Plant Molecular Biology Labfase (1993) by R.D.D. Croy, jointly published by BIOS Scientific Publications Ltd (UK) and Blackwell Scientific Publications (UK).

The invention is hereunder further explained by way of examples, including a material and method section, without being restrictive in the scope of the present invention.

Short description of the drawings:

Figure 1. Southern blots of the wild type (C24), and transgenic *CycB1;1* overexpressing *A. thaliana* lines (Cyc 28.10 and Cyc 5.9). Genomic DNA extracted from C24, Cyc 28.10 and Cyc 5.9 was digested with the indicated enzymes, separated on a 3% agarose gel and blotted. The membranes are hybridised with a probe derived from the CycB1;1 cDNA at high stringency.

Figure 2. RNA gel blot analysis. RNA was extracted from control plants (C24), homozygous Cyc 5.9 plants, homozygous Cyc 28.10 plants, and heterozygous Cyc 5.9 x CDC2aAt plants. 20 μg of RNA was separated on a 0.8% agarose gel and blotted on a nitro-cellulose membrane. Equal loading was confirmed by methylene blue staining. The blot was hybridised using an antisense CYCB1;1 probe.

Figure 3. CYCB1;1 (A) and CDC2aAt (B) western blots. Proteins were extracted from the indicated lines, and separated on a 12% SDS-PAGE gel. After immuno-blotting on a nitro-cellulose membrane the filters were probed using a CYCB1;1 (diluted 1/500) or a CDC2aAt (diluted 1/5000) specific antibody. As second antibody an anti-rabbit antibody coupled to peroxidase was used (diluted 1/10000). The detection was performed using the chemoluminescent procedure (Pierce, Rockford, IL).

Figure 4. Total root growth between day 2 and day 11. Genotypes are indicated as follows: c24, wild type; 5.9 and 28.10, independent CYCB1;1 overexpressing lines and cdc2a, 35S-CDC2a; the crosses "x" indicates female x male. Data indicate mean \pm SE (n = 8-13).

Figure 5. Root elongation rates as a function of time after sowing. Genotypes are indicated as in Fig 4. Data are averages \pm SE (n =8 -13) from the same roots as in Fig 4.

Figure 6. Length distribution of cortical cells along the roots of wild type (C24) and F1 seedlings of the cross between CYCB1;1 and CDC2a over expressing lines on day 9. Data are averages \pm SE (n = 2).

Figure 7. Area of the shoot in lateral projection at day 13 as determined from the image of the shoot on the culture plates. Genotypes are indicated as in Fig 4. Data are averages \pm SE (n =8 -13) from the same roots as in Fig 4.

The examples illustrate the invention without being intended as limiting:

EXAMPLES

General methodology for growth analysis in Arabidopsis thaliana.

Seeds of wild type *A. thaliana* (C24) and all transgenic lines are harvested from plants growing in the same conditions (22 °C and continuous light: 110 μ E m⁻² s⁻¹ PAR) and stored at 4°C after harvesting from completely dried plants. For each line to be tested the following screens were performed comparing wild-type and transgenic.

Screen 1. Root growth.

At day minus 3 (d–3) seeds were surface sterilised with 15% household bleach and plated on the surface of agar-solidified modified Hoagland solution (containing 4mM KNO₃, 1mM Ca (NO₃)₂, 2mM KH₂PO₄, 0.3mM MgSO₄, 0.09 μ M FeEDTA, 46.26 μ HBO₃, 9.91 μ M MnCl₂, 0.77 μ M ZnSO₄, 0.31 μ M CuSO₄, 0.11 μ M NaMoO₃, 0.1% (w/w) Sucrose and 0.8% (w/w) plant tissue culture agar (Lab M, Bury, England) in 12x12 cm square tissue culture plates, which were then placed vertically in the dark at 4°C for 3 days. On day 0, the plates transferred to a growth chamber with constant conditions (22 °C and 80 μ E m⁻² s⁻¹ PAR).

After germination, usually at day 2-3, the position of the tip of the main root was marked daily at the bottom of the plate with a razorblade. At day 11, the plates were digitised using a flatbed scanner with overhead illumination attached to a PC. Files with a resolution of 9.775 pixels per mm were saved in Tiff format.

Daily growth of individual roots on the plates was determined by measuring the distance along the main root axis between successive marks, using the freehand tool of the image analysis program Scion Image (Beta 3, Scion Corporation, Maryland, USA). The obtained data were transferred to the spreadsheet program Excel (Microsoft Corp). Total root growth over the observation period was obtained by adding all measured distances for each root. Root elongation rate for each root as a function of time was determined by dividing daily growth by the time interval between successive marks.

Estimation of the effect on cell division was done by measuring mature cell length. To this end the roots were whole mounted in the same nutrient solution specified above without the agar. Cortical cells were visualized with DIC optics in the region where root hairs have just reached their mature length using a microscope (Zeiss Axioscope fitted with a Zeiss Fluar 20X na = 0.75 lens). Typically, 5 average growing roots from the same experiment as used for determination of root elongation rate were selected. In each of these, the length of at least 20 mature cortical cells was measured on-line in the image analysis program Scion Image. For this, images were captured with a DCC camera (COHU 4910, USA) mounted on the microscope and connected to a PC with a framegrabber board (Scion LG3, Scion Corporation, Maryland, USA). The rate of cell production by the meristem was estimated from the ratio of the average root growth rate determined above, and mature cortical cell length.

In limited cases, cortical cell length was determined over the whole growth zone. Using the same setup as for the mature cell length, a series of partially overlapping images was recorded covering the whole of the growth zone and part of the mature region. Each series was transformed into a single composite image, which was then used to measure the length of all cells in each cortical file, starting from the quiescent centre. These data were transformed to express the length of each cell as a function of its midpoint. Interpolation and smoothing of these data was performed with a specially created algorithm, which repeatedly fits polynomials to small sections of the

data to estimate the midpoint of such section (Beemster and Baskin, 1998). Data obtained with this algorithm are equidistally spaced and were averaged between replicate roots. All data processing was done using the spreadsheet program Excel (Microsoft Corporation). This procedure gives, in addition to the mature cell length, also an idea about size of meristematic cells, and size of the meristem.

Screen 2: Total leaf area.

Total leaf (shoot) area was determined from the scanned images of the root systems described above. This was done by thresholding the image so as to select the entire shoot of each plantlet and then using the "analyse particles" routine. Obtained area values contain both leaf blade, petiole and hypocotyl. They are an underestimation for the true value of the area of these parts as the blades are aligned randomly instead of parallel to the field of view and there is overlap between various plant parts. It is obvious that the degree of underestimation increases for larger plants (more and larger organs) and therefore observed differences between genotypes are a conservative estimation of the true magnitude of differences in leaf area.

Screen 3: Leaf initiation and appearance rate.

At day 0 seeds were surface sterilised with 15% household bleach and plated on the surface of agar-solidified modified full strength Murashige and Skoog solution (Micro and macro elements (Duchefa)), 0.1% (w/w) Sucrose and 0.6% (w/w) plant tissue culture agar (Lab M, Bury, England) in 14.5 cm round / 2.5 cm high tissue culture plates (Falcon) which were placed horizontally in a growth chamber with constant conditions (22 °C and 110 µE m⁻² s⁻¹ PAR). After 7,14 and 21 days 5 average plants were selected from each line. On these plants the number of visible leaves was determined, and subsequently, by dissection under a binocular, the number of initiated leaves (= the number of visible leaves + the number of primordia).

Screen 4: Mature leaf size, cell density and estimated number f cells per leaf.

On the same plants from Screen 3, at day 21 all leaves were dissected and placed on a microscope slide in some water and covered with a coverslip. These slides were then scanned using the fladbed scanner (see Screen 1) and the area of each leaf was measured individually using the automated routine "analyse particles" on thresholded images in the aforementioned image analysis program Scion Image. After observation, the leaves were cleared overnight in ethanol and stored in 70% lactic acid. Cell density was determined by mounting the cleared leaves in lactic acid on a microscope slide and observing them with under DIC optics using the microscope fitted with DCC camera and connected to a PC with a framegrabber board (Scion LG3, Scion Corporation, Maryland, USA). On each leaf 4 images were captured of both adaxial epidermis and palisade parenchyma, halfway along the blade and in the middle between the mid vein and the leaf margin. On each of those images cell density was estimated by counting the number of cells in the image, whereby partial cells touching the left and upper edge were not, and those touching the right and lower edge of the image were included in the count. In the epidermis, stomata and pavement cells were counted separately. Cell density was then calculated as the number of cells divided by the area of the image. The density per blade was estimated as the average over the 4 images and the number of cells per blade of each type by multiplying this number with the total leaf area.

Screen 5: Seed weight and size.

Plants are germinated on agar as in Screen 1, transferred to soil at two weeks and all seeds are harvested when the plants have completely dried. For each plant the total seed weight is then determined. Finally seed size is determined by placing between 100 and 300 seeds per parental plant on the flatbedscanner. Images are scanned at 2400 dpi and analysed using the program Photoshop with a set of additional image analysis plug-ins (The

image processing toolkit version 3.0, Reindeer Games, Inc). The procedure is as follows: First the image is thresholded to select the seeds. Then touching seeds are separated using the watershed routine. After that all size/shape parameters are determined using the features/measure all command. From the resulting file the columns containing area, length, breadth, formfactor and roundness are selected. Outliers (dust and contamination particles) are removed based on their deviating formfactor and roundness factor. Of the remaining seeds the distribution is plotted and mean, median, average, standard deviation and standard error of the mean are determined.

EXAMPLE 1

Construction of the binary vector PGSC-TCyc1

Vector pcyc1T735 (gift of Dr. Paulo Ferreira, Departemento de Bioquímica Médica, UFRJ, Rio de Janeiro, Brazil), a PUC 19 vector (Yanish-Perron, 1985) containing 1.2Kb *CYCB1;1* cDNA with a T7 leader peptide and a NOS terminator, was digested with the restriction enzymes Ncol and Xbal. The resulting sequence consisting of 1.2Kb *CYCB1;1* cDNA, a T7 leader peptide and NOS terminator, was subsequently cloned into the vector TXGUS, (De Veylder et al 1998), from which the *GUS* gene had been excised with Ncol and Xbal, resulting in vector TXCyc1. TXCyc1 was digested with EcoRl and blunt cloned into the binary vector PGSC 1704, that carries hygromycine resistance, by means of the Snabl site, giving rise to vector PGSC-TCyc1.

EXAMPLE 2

Agrobacterium-mediated DNA transfer and plant transformation.

The PGSC TCyc1 vector was mobilised by the helper plasmid pRK 2013 into Agrobacterium tumefasciens C58C1RifR, harbouring the plasmid PGV 2260 (Deblaere, 1985). A. thaliana plants (ecotype C24) were then transformed by

44

root transformation (Valvekens, 1988). Transgenic plants were selected on hygromycine containing media and later transferred to soil for optimal seed production. A segregation analysis of ten independent lines was performed in the F1 generation based on hygromycine resistance, and out of two parental lines with single locus insertion (1/4 segregation; line 5 and 28) two homozygous daughter lines (5.9 and 28.10) of the F2 generation were selected. Southern blotting confirmed independency of the transformants (Fig. 1). To verify if lines 5.9 and 28.10 are actually overexpressing *CycB1;1*, a Northern (Fig. 2) and Western (Fig. 3) blots were performed using a *CYCB1;1* specific probe or antibody, respectively.

Both of these *CYCB1;1* overexpressing lines were crossed with a transgenic homozygous line of *A. thaliana*, containing the *CDC2a*At cDNA under control of a CaMV 35S promoter (Hemerly, 1995).

EXAMPLE 3

Series of reciprocal crosses

A series of reciprocal crosses (i.e. with each line functioning once as the pollen parent and once as seed parent) were made between the *CYCB1;1* overexpressing lines 5.9 and 28.10 and 35S-*CDC2a* and between these lines and the wild type. All seeds were harvested from a single batch of plants grown in a growth chamber with constant conditions (22 °C and 110 µE m⁻² s⁻¹ PAR), to avoid effects of the growth conditions of the mother plants on the progeny being analysed.

Root growth and cell length distribution was determined using <u>Screen 1</u> described above. Leaf (shoot) area was determined using <u>Screen 2</u> described above.

Total root growth

Overexpression of CYCB1;1 does not result in an increased root growth between day 2 and day 11 (Fig. 4), whereas overexpression of CDC2a in this

45

experiment even resulted in a 33% growth reduction. Interestingly, the F1 of the crosses between the *CYCB1;1* and *CDC2a* overexpressing lines grew 20 – 23% faster than the wild-type, with the exception of the 35S-*CDC2a* x 28.10 cross which grew 14% slower. For the 5.9 x 35S-*CDC2a* cross these results were consistent. Although not all combinations were tested, the similarity of the growth between the wild-type and the crosses of the overexpression lines with the wild-type indicate that for growth stimulation the combined overexpression of both *CDC2a* and *CYCB1;1* is required under these conditions.

Root elongation rates

In accordance with the absence of overall growth differences over the period between day 2 and day 11 between the wild type and the two *CYCB1;1* overexpressing lines (Fig 4), no significant difference in growth rate between these lines and the wild type are observed at any time during the observation period (Fig 5a). The reduced growth of the *CDC2a* overexpressing line appears to be associated with a reduced acceleration over time rather than a proportional difference throughout the growth period (Fig 5a). In contrast, the seedlings overexpressing both *CYCB1;1* and *CDC2a*, grow proportionally faster than the wild type throughout the growth period (Fig 5b). The reduced growth in the of the cross between *CDC2a* and *CYCB1;1* overexpressing lines with the 35S-*CDC2a* as maternal line seems to be due to a reduced acceleration compared to the wild type, because the growth rates shortly after germination are very similar (Fig 5b). This pattern is similar to that observed for the *CDC2a* overexpressing line (Fig 5a).

Cell length distribution

Variations in root elongation can be due to differences in cell expansion or cell division characteristics. To get a first indication about the cellular basis of the observed growth enhancement, the length distribution of cortical cells was analysed along the root. Typically small meristematic cells are found at the tip

of the root. In both wild type and roots from F1 seedlings from the *CYCB1;1* and *CDC2a* overexpressing lines this region is approximately 500 \Box m long (Fig 6). Next to the meristem, a region of rapidly growing cells is located between approximately 500 and 1750 μ m from the quiescent centre of the root for both genotypes. Basal to 1750 μ m, cells have reached their mature cell size. Given the small sample size of 2 roots per genotype and the large standard errors for the F1 line, it is uncertain if the observed differences in cell size are real.

Shoot growth

The area of the shoot in lateral projection mirrors the differences found in root growth (cf. Figs 3 and 6). Although only a rough measure for shoot area, these data indicate that the growth increase in the *CYCB1;1* and *CDC2a* overexpressing lines is not restricted to the roots, but also occurs in the aerial parts of the plant.

EXAMPLE 4

Weight of seeds

In addition to the effect of the overexpression of *CYCB1;1* and *CDC2A* on vegetative growth the following experiment is performed to investigate its generative growth. For this, seeds from the same reciprocal crosses are sown germinated on plates with agar solidified medium as specified for Example 3. Two weeks after germination, individual seedlings are planted individually in pots with moist potting mix and placed in a growth chamber with constant conditions (22 °C and 110 µE m⁻² s⁻¹ PAR). For each plant, individual seed pods are harvested when ripe, but before it opens in order to prevent seeds from getting lost. Counts are made for each plant of the number of seeds in a number of pods in predetermined positions. The total seed weight for each plant is weighed and the average seed weight is determined by weighing 100 seeds from each plant. Seed weight and size are also determined according to the methods described in Screen 5.

EXAMPLE 5

Leaf growth under natural light conditions

In order to establish that the observed growth enhancement is not dependent on the specific growth conditions utilised for Example 3, an experiment similar to Example 3 is performed, but with a day/night cycle of 8/16 hrs. In addition to this, leaf growth under natural light conditions on soil (natural light intensity is too high for root growth) is investigated. For this, seeds of the same crosses as used in Example 3 are sown directly into potting soil and placed in glasshouse. Leaf area are measured at 1 weekly intervals from 5 representative plants of each cross. For this the leaf blades are dissected and placed on a flatbed scanner, which make a digital image of the leaves. Total leaf area is determined for each plant by measuring the combined blade area's using the thresholding option of the image analysis program Scion Image.

Similarly, to examine the growth of plants under suboptimal conditions, coexpressing transgenic plants (e.g. Cdc2b and CycA) are grown on media with 0.5X NaCl – thus providing an environmental stress or more particularly a salt stress. The response of the transgenic plants to this stress condition are assessed using the various Screens described previously.

EXAMPLE 6

Yield Increase in Rice: Production of transgenic rice plants overexpressing cdc2Os-1 and cycOs2

1. Cloning of the cDNAs encoding cdc2Os-1 and cycOs2

The nucleotide sequences encoding cdc2Os-1 and cycOs2 have been published (Sauter et al., 1995). The corresponding cDNAs are cloned using the RT-PCR technique. The primer pair used to amplify the cdc2Os-1 is CCATGGAGCAGTACGAGAA (SEQ ID NO:1) for the 5' side and

CAGTGTCATTGTACCATCTCAA (SEQ ID NO:2) for the 3' side. The expected size of the amplification product is 891 bp. The template is total RNA isolated from a rice IR 52 cell suspension culture (Lee *et al.* 1989) that has been previously reversed transcribed as a bulk. The PCR conditions chosen to amplify this sequence are: 40 cycles of denaturation at 92° for 10 sec, annealing at 60° for 10 sec, extension at 72° for 60 sec.

Concerning the oligos for cycOs2, they match ATGGAGAACATGAGATCTGA (SEQ ID NO:3) for the 5'end and TTACAGTGCCACGCTCTTGAG (SEQ ID NO:4) for the 3'side of the sequence. The expected size of the amplification product is 1259 bp. The following PCR conditions are used: 45 cycles of denaturation at 92° for 10 sec, annealing at 53° for 10 sec, and extension at 72° for 90 sec.

The Pfu polymerase is used in both cases to generate blunt end fragments.

2. Subcloning of the maize ubiquitin promoter into a binary vector

The maize ubiquitin promoter from plasmid pAHC17 (Christensen, 1996) is excised as PstI fragment (made blunt ended with Pfu polymerase) and subcloned into the XbaI site (filled-in) of the binary vector pBIBHYG to give the vector pBHU.

3. Subcloning of the cdc2Os-1 and cyclinOs2 cDNAs into the pBHU binary vector

The cDNA of cdc2Os-1 and cyclinOs2 are cloned into the Sacl site (trimmed off) of pPHU to produce pBHU-cdc2 and pBHU-cyc2. The vectors are then introduced into an *Agrobacterium tumefaciens* octopine strain, via electroporation (McCormac, 1998).

4. Rice transformation via Agrobacterium

A. tumefaciens bearing either pBHU-cdc2 or pBHU-cyc2 are used to produce transgenic rice expressing either cdc2Os-1 or cyclinOs2 under control of the ubiquitin promoter, following Hiei Y., (1994). Lines expressing highest levels

49

PCT/EP00/02441

of the transgenes are crossed to produce transgenic lines co-expressing both transgenes.

5. Expected results

On the basis of results obtained in *Arabidopsis* it is anticipated that transgenic rice plants overproducing both cdc2Os-1 and cyclinOs2 will display increased growth rates and robustness. Since the growth stimulating effects observed in *Arabidopsis* have rather general character and not confined to a particular organ or tissue, we expect also the transgenic rice to show an increase in the grain size.

EXAMPLE 7

Mutant alleles and wild type genes of CDC2bAt and CDC2fAt

Mutant alleles and wild type genes of CDC2bAt and CDC2fAt were cloned under the control of the CaMV 35S promoter and transferred to a binary vector. The CDC2bAt and CDC2fAt genes are cloned in a kanamycin-containing vector. The mutant alleles include dominant negative forms of CDKs (CDC2b.N161 and CDC2fAt.N164 - in both constructs the D residue was replaced with an N residue; this mutation has been shown to inactivate the kinase causing an arrest of the cell cycle) and positive forms of the CDKs. The following cyclin genes are cloned in a hygromycin vector allowing the coexpression of the cyclins and CDK combinations by crossing.

1. Plasmid constructions

CDC2bAt constructs:

Ncol and BamHI restriction sites were introduced in the cDNAs of CDC2bAt, CDC2bAt.A14F15 and CDC2bAt.N161 by performing PCRs with the following primers: 5'-GGCCATGGAGAAGTACGAGAAGC-3' (SEQ ID NO:5) (containing a Ncol restriction site) and 5'-GGGGATCCTCAGAACTGAGACTTGTCAAGG-3' (SEQ ID NO:6) (containing a BamHI restriction site). The different PCR fragments were cut with Ncol and

WO 00/56905 PCT/EP00/02441

BamHI and cloned into the Ncol and BamHI sites of pH35S (Hemerly et al., 1995). The cassettes 35S-CDC2bAt-3'NOS, 35S-CDC2bAt.A14F15-3'NOS and 35S-CDC2bAt.N161-3'NOS were cloned into the EcoRI and Sall restriction sites of the binary vector pBinPlus (Engelen, 1995 Transgenic Research, 4:288-290)

CDC2fAt constructs:

Clal and Sall restriction sites were introduced in the cDNAs of CDC2fAt, CDC2fAt.A26F27 and CDC2fAt.N164 by performing PCRs with the following primers: 5'-GGATCGATATGGACGAGGGAGTTATAGC-3' (SEQ ID NO.7) (containing a Clal restriction site) and 5'-GGGGGAAGCACAGTCGACATATGC-3' (SEQ ID NO:8) (containing a Sall restriction site. The different PCR fragments were cut with Clal and Sall and cloned into the Clal and Sall sites of the binary vector pGV1990.

CYCA2;2 constructs:

in order to introduce the Clal and Sall restriction sites and a HA tag into the cDNAs of CYCA2;2 and CYCA2;2 \(\Delta 64\), PCRs were performed using the following primers:

5'-GCGATCGATATGGGCTACCCTTACGATGTTCCAGATTACGCTATGTA
TTGCTCTTCTCGATGC-3' (SEQ ID NO:9) (containing a Clal site and a HA
tag in fusion with the full length *CYCA2;2*),

5'-GCGATCGATATGGGCTACCCTTACGATGTTCCAGATTACGCTACCTCT GCAGATATTATTTATTC-3' (SEQ ID NO:10) (containing a Clal site and a HA tag in fusion with the truncated CYCA2;2\u03b164) and

5'-GGCGTCGACGTTGCTTGGTGTCATCTTG-3' (SEQ ID NO: 11) (containing a Sall restriction site).

The different PCR fragments obtained were cut with Clal and Sall and cloned into the Clal and Sall sites of pGV1990.

CYCA2:3 constructs:

in order to introduce the Sall restriction sites and a HA tag into the cDNAs of CYCA2;3 and CYCA2;3\(\triangle 63\), PCRs were performed using the following primers:

5'-

GCGGTCGACATGGGCTACCCTTACGATGTTCCAGATTACGCTATGGGG AAGGAAAATGCTG-3' (SEQ ID NO:12) (containing a Sall site and a HA tag in fusion with the full length *CYCA2;*3),

5'-

GCGGTCGACATGGGCTACCCTTACGATGTTCCAGATTACGCTGTTAACT CCAATACAGC-3' (SEQ ID NO:13) (containing a Sall site and a HA tag in fusion with the truncated CYCA2;3\(\Delta\)63) and

5'- GGCGTCGACGGTTAGGAGTTGAAACC-3' (SEQ ID NO:14) (containing a Sall restriction site).

The 3'NOS was removed from pH35S by BamHI and XbaI digestion and cloned into the BamHI and XbaI restriction sites of the vector pLBR19, resulting into the pLBR19/NOS vector. The obtained PCR fragments were cut with SaII and cloned into the SaII restriction site of pLBR19/NOS. By restriction with the KpnI site, the cassettes 35S-CYCA2;3-3'NOS and 35S-CYCA2;3\(\Delta\)63-3'NOS were removed from pLBR19/NOS, blunt ended and cloned into the SNABI restriction site of the pGSC1704 binary vector.

CYCB2;1 constructs:

WO 00/56905 PCT/EP00/02441

52

in order to introduce the Sall and BamHl restriction sites and a HA tag into the cDNAs of CYCB2;1 and CYCB2;1 \(\Delta 44\), PCRs were be performed using the following primers:

5'-

GCGGTCGACATGGGCTACCCTTACGATGTTCCAGATTACGCTATGGTTA ACTCATGCGAG-3' (SEQ ID NO: 15) (containing a Sall site and a HA tag in fusion with the full length *CYCB2;1*),

5'-

GCGGTCGACATGGGCTACCCTTACGATGTTCCAGATTACGCTCAGAATC TCGCTGGTGC-3' (SEQ ID NO:16) (containing a Sall site and a HA tag in fusion with the truncated *CYCB2;1*\(\textit{\Delta}44\)) and

5'- CCGGATCCTGTACAAACTACTTAC-3' (SEQ ID NO: 17) (containing a BamHI restriction site).

The obtained PCR fragments were cut with Sall and BamHI and cloned into the Sall and BamHI restriction sites of pLBR19/NOS. By restriction with the KpnI site, the cassettes 35S-CYCB2;1-3'NOS and 35S-CYCB2;1\(\Delta 44-3'NOS\) were removed from pLBR19/NOS, blunt ended and cloned into the SNABI restriction site of the pGSC1704 binary vector.

CYCB-type∆ construct:

in order to introduce the Ncol and BamHI restriction sites and a HA tag into the cDNA of CYCB-type \(\Delta \), a PCR was performed using the following primers:

5'-

GCGCCATGGGCTACCCTTACGATGTTCCAGATTACGCTCCACATATCCG TGATGAGG-3' (SEQ ID NO:18) (containing a Ncol site and a HA tag in fusion with the truncated CYCB-type△), and 5'- GCGGATCCATTCTCCCCATTTTGG-3' (SEQ ID NO:19) (containing a BamHl restriction site).

The PCR fragment was cut with Ncol and BamHI and cloned into the Ncol and BamHI sites of pH35S. The cassette 35S-CYCB-type⊿-3'NOS were removed from pH35S by EcoRI and BamHI digestion, blunt ended and cloned into the SNABI restriction site of the pGSC1704 binary vector.

2. Plant transformation

The different constructs were transferred into *Agrobacterium tumefaciens* and introduced into *Arabidopsis thaliana* ecotype C24 by the floral dip method (Clough, 1998).

3. Analysis

Transgenic plants are analysis for various growth and cell division characteristics or phenotypes according to the Screens described in the General Methodology above.

EXAMPLE 8

Evidence of cell cycle protein complexes

1. Purification protocol of native CDK complexes

All steps are carried out at 4°C or on ice; at all stages the presence of CDK complexes is followed based on their kinase activity or via western analysis. Cell suspension cells and buffers:

five fold diluted homogenization buffer: the concentration of all additives is five times lower than in homogenization buffer except for Tris and NaCl which are the same (respectively 25 and 85 mM)

DEAE buffer: 50mM Tris.HCl (pH= 7.8 or 9.3) plus 5mM MgCl₂, 5mM EGTA, 5mM β-glycerophosphate, 1mM NaF, 0.01% NP40, 1mM DDT.

54

0.25mM PMSF, $1\mu g/ml$ aprotonin and leupeptin, 0.1mM benzamidine and NaVO₄.

S200 buffer: 50mM Tris.HCl (pH= 7.8 or 9.3) plus 15mM MgCl₂, 5mM EGTA, 5mM β -glycerophosphate, 1mM NaF, 1mM DDT, 0.1mM NaVO₄, 100mM NaCl.

Total extract: in homogenization buffer (85mM NaCl), typically 70ml **Cleansing step**: DEAE Sepharose FF (Pharmacia): flow through collection into 150ml superloop (including washing of the column in five fold diluted homogenization buffer) flow 10ml /min.

Affinity binding onto Cks homologues coupled to an affinity matrix:

The columns are connected in series in this specific order: P9 (Ckshs1) 10ml, P13 (Suc1) 3ml and P10 (Cks1At) 10ml. The sample (150ml) is loaded at a flow rate of 200µl/min.

The columns (still connected in series) are then washed with bead buffer to minimize non-specific interactions (cf. higher salt concentration) at a flow rate of 500µl/min.

In this way all CDC2a binds to the P9 column. CDC2b binds to both P13 and P10.

CDK- complex elution via excess of free ligand:

Affinity columns are disconnected from one another and eluted individually with an excess of their respective free ligand (applied in DEAE buffer, pH=7.8).

Preparation of an excess of free ligand is done as follows: lyophilized powder is dissolved in 6M urea in 25mM Tris pH=7.5 as to obtain a highly concentrated solution (typically 25mg/ml); then desalted on a Sephadex G25 column (1x10cm) to DEAE buffer (PH=7.8) and slowly injected onto the affinity columns in reversed flow mode at a flow rate of 200µl/min, thus displacing the bound proteins complexes which are collected.

2. Separation of affinity eluted CDK-complexes:

The CDK containing fractions are first concentrated on DEAE 650S (TSK) therefore the pH is raised to 9.3 and the sample is applied onto the column (HR5/5 Pharmacia). Bound proteins are eluted in one step through injection of 0.5M NaCL in DEAE buffer (pH 7.8).

Then the concentrated CDK fraction is further separated by size on a gel filtration column: Superdex 200pg Pharmacia (1.7x100cm column Omnifit) or a Sephacryl S200 Pharmacia (1.5x100cm column Pharmacia). The columns are equilibrated in S200 buffer when fractions (5ml) are collected or in DEAE buffer when the eluting proteins are immediately eluted onto DEAE 650S (TSK) (HR5/5 columns), in the latter case the pH of the size exclusion buffer was raised to 9.3 (cf. concentration step).

Separately collected CDK containing fractions (5ml) which are later bound on DEAE or those CDK complexes which are directly bound to DEAE when eluting from the size exclusion column are eluted in a similar way: a 10 column-volume gradient of 0–500mM NaCl is applied and the eluting complexes are collected.

A final purification step consists of hydrophobic interaction chromatography: The conductivity of the purified fractions is raised to100mS by adding saturated ammonium sulphate and the samples are individually applied onto Ether PW-5 or Phenyl PW-5 (TSK). The bound complexes are eluted with a decreasing (NH₄)₂SO₄ gradient and can be tested on their kinase activity.

3. Protein complexes purified

In the CDC2a fraction (eluted from CKShs1) there were two 100kDa combinations: CDC2a with CycB2;2 and with CycA2;2. Higher molecular weight complexes containing CDC2a were also detected.

WO 00/56905 PCT/EP00/02441 56

In the CDC2b fraction obtained from CKS1At both CycB1;1 and cyclin A2;2 were detected in 100kDa complexes, suggesting their association with CDC2b. CycB1;1 was also detected in a 200kDa complex and a higher molecular weight complex.

Some complexes are not bound during the affinity purification.

Cyclin A2:1 is present in a 100kDa complex with an as yet unidentified protein(s). However it is not yet clear whether the latter one is a CDC homologue or an (un)related protein.

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WO 00/56905 PCT/EP00/02441

61

explants by using kanamycine selection. Proc. Natl. Acad. Sci. USA 85: 5536-5540

TABLE 1 – Examples of cyclins in plants

Class of	Tvoical	Examples	Original	Comments	Reference
cyclin	phase		name		
	dependence				
A 1	S/G2/M	Zeama;CYCA1;1	cyclIZm	Zeama;CYCA1;1 triggers frog	see Renaudin et al (1996) Plant
		Nicta;CYCA1;1	ntcyc25	oocyte maturation;	Mol. Biol. 32: 1003-1018
	-			Nicta;CYCA1;1 rescues G1	
				cyclin deficiency in budding	
A2	S/G2/M	Nicta;CYCA2;1	ntcyc27	Medsa;CYCA2;1 expression	see Renaudin et al (1996) Plant
		Medsa;CYCA2;1 ^c	cycMs3	suppresses the	Mol. Biol. 32: 1003-1018; Day
				α-pheromone-induced cell cycle	and Reddy (1998) Plant Mol.
				arrest in yeast. Medsa;CYCA2;1	Biol 36: 451-461
				and Nicta;CYCA2;1	
				complement G1 cyclin	
				deficiency in yeast	
A3	S/early G2	Catro;CYCA3;1	CYS	Catro;CYCA3;1 rescues G1	see Renaudin et al (1996) Plant
				cyclin deficiency in yeast	Mol. Biol. 32: 1003-1018
81	G2/M	Arath;CYCB1;1	cyc1At	Arath;CYCB1;1,	see Renaudin et al (1996) Plant
		Arath;CYCB1;2	cyc1bAt	Zeama;CYCB1;1,	Mol. Biol. 32: 1003-1018
		Catro;CYCB1;1	CYM	Zeama;CYCB1;2, and	
		Nicta;CYCB1;2	NycycZ9	Glyma;CYCB1,2 trigger frog	
		Zeama;CYCB1;1	cyclaZm	oocyte maturation.	
		Zeama;CYCB1;2	cyclbZm	Arath;CYCB1;2,	
		Glyma;CYCB1;1	S13-6	Catro;CYCB1,1, and	
				Nicta;CYCB1;2 rescue G1	
				cyclin deficiency in yeast	
B2	G2/M	Arath;CYCB2;2	cyc2bAt	Zeama;CYCB2;1 triggers	Hemerly et al (1994) PNAS
		Zeama;CYCB2;1	cycllIZm	oocyte maturation;	91:11313-11317; Renaudin et al
		Medsa;CYCB2;2	cycMs2	Medsa;CYCB2;2-immunoprecipi	(1994) PNAS 91: 7375-7379;

		Orysa; CYCB2;1 Orysa;CYCB2;2		tated kinase activity is maximal in G2	Meskiene et al (1995) Plant Cell 7: 759-771;
DJ	Unknown	Arath;CYCD1;1 Antma; CYCD1;1 Heltu;CYCD1;1	cyclin 81	Rescues G1 deficiency in yeast; associates with CDC2aAt in the two-hybrid system	Soni et al (1995) Plant Cell 7:1847-1857; Antma and Heltu - unpublished data see: Sorrell et al (1999) Plant Physiol. 119:
D2	Non specific	Arath;CYCD2;1 Nicta;CYCD2;1 Cheru;CYCD2;1	cyclin 82	Rescues G1 deficiency in yeast Expression sucrose inducible; Nicta;CYCD2;1 transcript peaks during M	Sorrell et al (1999) Plant Physiol. 119: 343-351; Soni et al (1995) Plant Cell 7:1847- 1857; Renz et al (1997) Plant Physiol. 113: 1004.
D3	Non specific	Arath;CYCD3;1 Medsa;CYCD3;1 Nicta;CYCD3;1 Nicta;CYCD3;2 Antma;CYCD3;2 Antma;CYCD3;1 Heltu;CYCD3;1	cyclin53 cycMs4	Rescues G1 deficiency in yeast. Expression cytokinin inducible; Nicta;CYCD3;1 transcript peaks during M. Expressed in only a subset of proliferating cells; interacts with Rb and ICK1	Riou-Khamlichi et al (1999) Science 283: 1541-1544 (Antma, A. majus; Heltu, Helianthus tuberosus – unpublished data see: Sorrell et al (1999) Plant Physiol. 119:
D4	Unknown	Arath;CYCD4;1		Expression sucrose inducible; expressed during lateral root primordia formation	De Veylder et al 1999, Plant, 208: 453-462

a At the transcriptional level. b Nomenclature according to Renaudin et al. (1996). c Expressed in a non-specific manner.

TABLE 3

EXEMPLARY TISSUE SPECIFIC FOR USE IN THE PERFORMANCE OF THE PRESENT INVENTION

GENE SOURCE	EXPRESSION PATTERN	REFERENCE
α-amylase (<i>Amy32b</i>)	aleurone	Lanahan, M.B., e t al., Plant Cell 4:203-211, 1992; Skriver, K., et al. Proc. Natl. Acad. Sci. (USA) 88: 7266-7270, 1991
cathepsin β-like gene	aleurone	Cejudo, F.J., et al. Plant Molecular Biology 20:849-856, 1992.
Agrobacterium rhizogenes rolB	cambium	Nilsson et al., Physiol. Plant. 100:456-462, 1997
PRP genes	cell wall	http://salus.medium.edu/mmg/tiern ey/html
AtPRP4	flowers	http://salus.medium.edu/mmg/tiern ey/html
chalene synthase (chsA)	flowers	Van der Meer, et al., Plant Mol. Biol. 15, 95-109, 1990.
LAT52	anther	Twell et al Mol. Gen Genet. 217:240-245 (1989)
apetala-3	flowers	
chitinase	fruit (berries, grapes, etc)	Thomas et al. CSIRO Plant Industry, Urrbrae, South Australia, Australia; http://winetitles.com.au/gwrdc/csh9 5-1.html
rbcs-3A	green tissue (eg leaf)	Lam, E. et al., The Plant Cell 2: 857-866, 1990.; Tucker et al., Plant Physiol. 113: 1303-1308, 1992.
leaf-specific genes	leaf	Baszczynski, et al., Nucl. Acid Res. 16: 4732, 1988.
AtPRP4	leaf	http://salus.medium.edu/mmg/tiern ey/html
Pinus cab-6	leaf	Yamamoto et al., Plant Cell Physiol. 35:773-778, 1994.
SAM22	senescent leaf	Crowell, et al., Plant Mol. Biol. 18: 459-466, 1992.
R. japonicum nif gene	nodule .	United States Patent No. 4, 803, 165
B. japonicum nifḤ gene	nodule	United States Patent No. 5, 008, 194
GmENOD40	nodule	Yang, et al., The Plant J. 3: 573-585.
PEP carboxylase (PEPC)	nodule	Pathirana, et al., Plant Mol. Biol. 20: 437-450, 1992.
leghaemoglobin (Lb)	nodule	Gordon, et al., J. Exp. Bot. 44:

	<u> </u>	1453-1465, 1993.
Tunana haaillifarm virus aana	nhloom.	<u> </u>
Tungro bacilliform virus gene	phloem	Bhattacharyya-Pakrasi, et al, The Plant J. 4: 71-79, 1992.
sucrose-binding protein gene	plasma membrane	Grimes, et al., The Plant Cell 4:1561-1574, 1992.
pollen-specific genes	polien; microspore	Albani, et al., Plant Mol. Biol. 15: 605, 1990; Albani, et al., Plant Mol. Biol. 16: 501, 1991)
Zm13	pollen	Guerrero et al Mol. Gen. Genet. 224:161-168 (1993)
apg gene	microspore	Twell et al Sex. Plant Reprod. 6:217-224 (1993)
maize pollen-specific gene	pollen	Hamilton, et al., Plant Mol. Biol. 18: 211-218, 1992.
sunflower pollen-expressed gene	pollen	Baltz, et al., The Plant J. 2: 713-721, 1992.
B. napus pollen-specific gene	pollen;anther; tapetum	Arnoldo, et al., J. Cell. Biochem., Abstract No. Y101, 204, 1992.
root-expressible genes	roots	Tingey, et al., EMBO J. 6: 1, 1987.
tobacco auxin-inducible gene	root tip	Van der Zaal, et al., Plant Mol. Biol. 16, 983, 1991.
β-tubulin	root	Oppenheimer, et al., Gene 63: 87, 1988.
tobacco root-specific genes	root	Conkling, et al., Plant Physiol. 93: 1203, 1990.
B. napus G1-3b gene	root	United States Patent No. 5, 401, 836
SbPRP1	roots	Suzuki et al., Plant Mol. Biol. 21: 109-119, 1993.
AtPRP1; AtPRP3	roots; root hairs	http://salus.medium.edu/mmg/tiern ey/html
RD2 gene	root cortex	http://www2.cnsu.edu/ncsu/researc
TobRB7 gene	root vasculature	http://www2.cnsu.edu/ncsu/researc
AtPRP4	leaves; flowers; lateral root primordia	http://salus.medium.edu/mmg/tiern ey/html
seed-specific genes	seed	Simon, et al., Plant Mol. Biol. 5: 191, 1985; Scofield, et al., J. Biol. Chem. 262: 12202, 1987.; Baszczynski, et al., Plant Mol. Biol. 14: 633, 1990.
Brazil Nut albumin	seed	Pearson, et al., Plant Mol. Biol. 18: 235-245, 1992.
legumin	seed	Ellis, et al., Plant Mol. Biol. 10: 203- 214, 1988.
glutelin (rice)	seed	Takaiwa, et al., Mol. Gen. Genet. 208: 15-22, 1986; Takaiwa, et al., FEBS Letts. 221: 43-47, 1987.
zein	seed	Matzke et al Plant Mol Biol, 14(3):323-32 1990

парА	seed	Stalberg, et al, Planta 199: 515- 519, 1996.
wheat LMW and HMW glutenin-1	endosperm	Mol Gen Genet 216:81-90, 1989; NAR 17:461-2, 1989
wheat SPA	seed	Albani et al, Plant Cell, 9: 171-184, 1997
wheat α, β, γ-gliadins	endosperm	EMBO 3:1409-15, 1984
barley Itr1 promoter	endosperm	
barley B1, C, D, hordein	endosperm	Theor Appl Gen 98:1253-62, 1999; Plant J 4:343-55, 1993; Mol Gen Genet 250:750-60, 1996
barley DOF	endosperm	Mena et al, The Plant Journal, 116(1): 53-62, 1998
blz2	endosperm	EP99106056.7
synthetic promoter	endosperm	Vicente-Carbajosa et al., Plant J. 13: 629-640, 1998.
rice prolamin NRP33	endosperm	Wu et al, Plant Cell Physiology 39(8) 885-889, 1998
rice α-globulin Glb-1	endosperm	Wu <i>et al,</i> Plant Cell Physiology 39(8) 885-889, 1998
rice OSH1	embryo	Sato et al, Proc. Natl. Acad. Sci. USA, 93: 8117-8122, 1996
rice α-globulin REB/OHP-1	endosperm	Nakase et al. Plant Mol. Biol. 33: 513-522, 1997
rice ADP-glucose PP	endosperm	Trans Res 6:157-68, 1997
maize ESR gene family	endosperm	Plant J 12:235-46, 1997
sorgum α-kafirin	endosperm	PMB 32:1029-35, 1996
KNOX	embryo	Postma-Haarsma et al, Plant Mol. Biol. 39:257-71, 1999
rice oleosin	embryo and aleuron	Wu et at, J. Biochem., 123:386, 1998
sunflower oleosin	seed (embryo and dry seed)	Cummins, et al., Plant Mol. Biol. 19: 873-876, 1992
LEAFY	shoot meristem	Weigel <i>et al., Cell 6</i> 9:843-859, 1992.
Arabidopsis thaliana knat1	shoot meristem	Accession number AJ131822
Malus domestica kn1	shoot meristem	Accession number Z71981
CLAVATA1	shoot meristem	Accession number AF049870
stigma-specific genes	stigma	Nasrallah, et al., Proc. Natl. Acad. Sci. USA 85: 5551, 1988; Trick, et al., Plant Mol. Biol. 15: 203, 1990.
class I patatin gene	tuber	Liu et al., Plant Mol. Biol. 153:386- 395, 1991.
PCNA rice	meristem	Kosugi <i>et al, Nucleic Acids Research 19:</i> 1571-1576, 1991; Kosugi S. and Ohashi Y, <i>Plant Cell</i> 9:1607-1619, 1997.
Pea TubA1 tubulin	Dividing cells	Stotz and Long, <i>Plant Mol.Biol.</i> 41, 601-614. 1999
Arabidopsis cdc2a	cycling cells	Chung and Parish, FEBS Lett,

		3;362(2):215-9, 1995
Arabidopsis Rop1A	Anthers; mature pollen + pollen tubes	Li et al. 1998 Plant Physiol 118, 407-417.
Arabidopsis AtDMC1	Meiosis-associated	Klimyuk and Jones 1997 <i>Plant J.</i> 11, 1-14.
Pea PS-IAA4/5 and PS-IAA6	Auxin-inducible	Wong et al. 1996 Plant J. 9, 587-599.
Pea farnesyltransferase	Meristematic tissues; phloem near growing tissues; light- and sugar-repressed	Zhou et al. 1997 <i>Plant J</i> . 12, 921- 930
Tobacco (N. sylvestris) cyclin B1;1	Dividing cells / meristematic tissue	Trehin et al. 1997 Plant Mol.Biol. 35, 667-672.
Catharanthus roseus Mitotic cyclins CYS (A-type) and CYM (B-type)	Dividing cells / meristematic tissue	Ito et al. 1997 Plant J. 11, 983-992
Arabidopsis cyc1At (=cyc B1;1) and cyc3aAt (A-type)	Dividing cells / meristematic tissue	Shaul et al. 1996 Proc.Natl.Acad.Sci.U.S.A 93, 4868-4872.
Arabidopsis tef1 promoter box	Dividing cells / meristematic tissue	Regad et al. 1995 Mol.Gen.Genet. 248, 703-711.
Catharanthus roseus cyc07	Dividing cells / meristematic tissue	Ito et al. 1994 Plant Mol.Biol. 24, 863-878.

TABLE 4 EXEMPLARY CONSTITUTIVE PROMOTERS FOR USE IN THE PERFORMANCE OF THE PRESENT INVENTION

GENE SOURCE	EXPRESSION PATTERN	REFERENCE
Actin	constitutive	McElroy et al, Plant Cell, 2: 163- 171, 1990
CAMV 35S	constitutive	Odell <i>et al,</i> Nature, 313: 810-812, 1985
CaMV 19S	constitutive	Nilsson et al., Physiol. Plant. 100:456-462, 1997
GOS2	constitutive	de Pater <i>et al,</i> Plant J Nov;2(6):837-44, 1992
ubiquitin	constitutive	Christensen <i>et al</i> , Plant Mol. Biol. 18: 675-689, 1992
rice cyclophilin	constitutive	Buchholz et al, Plant Mol Biol. 25(5): 837-43, 1994
maize H3 histone	constitutive	Lepetit et al, Mol. Gen. Genet. 231:276-285, 1992
actin 2	constitutive	An <i>et al,</i> Plant J. 10(1); 107-121, 1996

CLAIMS

- 1. A method for modifying plant growth and/or yield or modifying architecture comprising introducing into a plant cell, plant tissue or plant (a) nucleic acid molecule(s) or regulatory sequence(s), wherein the introduction of the nucleic acid molecule(s) or regulatory sequence(s) result(s) in an increased or de novo expression of at least two cell cycle interacting proteins capable of forming a (heteromeric) complex in a plant cell.
- 2. The method of claim 1, wherein said nucleic acid molecule(s) encode(s) said cell cycle interacting protein(s) and the regulatory sequence(s) is (are) capable of increasing the expression of a gene encoding said cell cycle interacting protein(s).
- 3. The method of claim 1 or 2, wherein one of said cell cycle interacting proteins is a protein kinase.
- 4. The method of any one of claims 1 to 3, wherein said protein kinase is a cycline-dependent kinase (CDK).
- 5. The method of claim 4, wherein said CDK is an A-type or a B-type CDK.
- 6. The method of claim 4 or 5, wherein said CDK is a PSTAIRE type CDK.
- 7. The method of claim 5, wherein the A-type CDK is Cdc2a and B-type CDK is Cdc2b.
- 8. The method of any one of claims 1 to 7, wherein one of said cell cycle interactiving proteins is a cyclin (Cyc).

- 9. The method of claim 8, wherein the cyclin is a G1 cyclin.
- 10. The method of claim 8 or 9, wherein the cyclin is A, B, C, D or E-type cyclin.
- 11. The method of any one of claims 8 to 10, wherein the cyclin is a CycA1;1, CycA2;1, CycA2;2, CycA2;3, CycA3;1, CycB1;1, CycB1;2, CycB2;1, CycB2;2, CycD1;1, CycD2;1, CycD3;1 or CycD4;1.
- 12. The method of claim 11, wherein the cyclin is CycD4;1.
- 13. The method of any one of claims 1 to 12, wherein one of the cell cycle interacting proteins is a ORC1, CDC6, CDC7, DBF4, E2F or DP.
- 14. The method of claim 13, wherein
 - (a) one cell cycle interacting protein is ORC1 and one is CDC 6;
 - (b) one cell cycle interacting protein is DBF4 and one is CDC 7;
 - (c) one cell cycle interacting protein is E2F and one is DP.
- 15. The method of any one of claims 1 to 12, wherein one cell cycle interactiving protein is a CDK and one of said cell cycle interactive proteins is a cyclin.
- 16. The method of claim 15, wherein
 - (a) the CDK is a A-type CDK and the cyclin is a B-type cyclin;
 - (b) the CDK is a B-type CDK and the cyclin is a B-type cyclin;
 - (c) the CDK is a A-type CDK and the cyclin is a D-type cyclin; or
 - (d) the CDK is a B-type CDK and the cyclin is a A-type cyclin.
- 17. The method of claim 15 or 16, wherein
 - (a) the cyclin is CycA2;1 and the CDK is Cdc2a;
 - (b) the cyclin is CycA2;2 and the CDK is Cdc2a, Cdc2b, Cdc2f, Cdc2bN161 or Cdc2aN146;

- (c) the cyclin is CycA2;3 and the CDK is Cdc2b, Cdc2f, Cdc2bN161 or Cdc2aN146;
- (d) the cyclin is CycB1;1 and the CDK is Cdc2a or Cdc2b;
- (e) the cyclin is CycB1;2 and the CDK is Cdc2a or Cdc2b;
- (f) the cyclin is CycB2;1 and the CDK is Cdc2b or Cdc2f;
- (g) the cyclin is CycB2;2 and the CDK is Cdc2a or Cdc2b;
- (h) the cyclin is CycD1;1 and the CDK is G1-CDK or Cdc2a;
- (i) the cyclin is CycD2;1 and the CDK is G1-CDK or Cdc2a;
- (j) the cyclin is CycD3;1 and the CDK is G1-CDK or Cdc2a; or
- (k) the cyclin is CycD4;1 and the CDK is Cdc2a, Cdc2b, Cdc2f, Cdc2bN161, Cdc2aN146 or Cdc2fN164.
- 18. The method of any one of claims 1 to 17 wherein the CDK is Cdc2a and the cyclin is CycD4;1.
- 19. The method of anyone of claims 1 to 18, wherein the cell cycle interacting protein is a modified, a homolog or an analog form of a cell cycle interacting protein.
- 20. The method of any one of claims 1 to 19, wherein said nucleic acid molecule(s) encode(s) at least a catalytic and/or regulatory subunit of said cell cycle interacting protein(s).
- 21. The method of any one of claims 1 to 20, wherein said cell cycle interacting proteins are expressed in one or more particular plant cells, tissues, organs and plant parts and progeny plants.
- 22. The method of any one of claims 1 to 21, wherein said nucleic acid molecule(s) is (are) operatively linked to regulatory sequences allowing the expression of the nucleic acid molecule(s) in the plant cell.

- 23. The method of any one of claims 1 to 22, wherein the regulatory sequence comprises promoter, enhancer, silencer, intron sequences, 3'UTR and/or 5'UTR regions, protein and/or RNA stabilizing elements.
- 24. The method of any one of claims 1 to 23, wherein said regulatory sequence is a chimeric, tissue specific, constitutive or inducible promotor.
- 25. The method of any one of claims 1 to 24, wherein said plant is a monocotyledonous or a dicotyledonous plant.
- 26. The method of any one of claims 1 to 25 wherein said plant is a crop plant, root plant, oil producing plant, wood producing plant, agricultured bioticultured plant, fodder or forage legume, companion plant or horticultured plant.
- 27. The method of claim 25 or 26, wherein said plant is wheat, barley, maize, rice, carrot, sugar beet, cichorei, cotton, sunflower, tomato, cassava, grapes, soybean, sugar cane, flax, oilseed rape, tea, canola, onion, asparagus, carrot, celery, cabbage, lentil, broccoli, cauliflower, brussel sprout, artichoke, okra, squash, kale, collard greens or potato.
- 28. A nucleic acid molecule encoding at least two cell cycle interacting proteins as defined in any one of claims 1 to 21.
- 29. A vector comprising the nucleic acid molecule of claim 28 or at least two nucleic acid molecules and/or regulatory sequences as defined in any one of claims 1 to 21.
- 30. The vector of claim 29 comprising separate nucleic acid molecules encoding at least one of said cell cycle interacting proteins as defined in any one of claims 1 to 21.

- 31. A composition comprising vectors wherein each vector contains at least one nucleic acid molecule encoding at least one cell cycle interacting protein as defined in any one of claims 1 to 21; and wherein the expression of said vectors results in the production of at least two cell cycle interacting proteins and assembly of the same in a complex in vitro or in vivo.
- 32. The vector of claim 29 or 30 or the composition of claim 31 wherein the nucleic acid molecule is operatively linked to (a) control sequence(s) allowing the expression of cell cycle interacting proteins in a host cell.
- 33. The vector or composition of claim 32 wherein said control sequence is a constitutive, chimeric, tissue specific or inducible promoter.
- 34. A host cell comprising the nucleic acid molecule of claim 28, the vector of any one of claims 29, 30, 32 or 33, or the composition of claims 31 or 32.
- 35. A method for the preparation of a cell cycle protein complex comprising:
 - (a) culturing the host cell of claim 34 under conditions suitable for the expression of the nucleic acid molecules; and
 - (b) recovering the complex from the culture.
- 36. A cell cycle protein complex obtainable by the method of claim 35 or encodable by the nucleic acid molecule of claim 28.
- 37. A transgenic plant cell displaying an increased amount of or de novo cell cycle interacting protein complex compared to a corresponding wild type plant cell said transgenic plant cell comprising at least one nucleic acid molecule or regulatory sequence as defined in any one of claim 1 to 21, a nucleic acid molecule of claim 28, the vector of any one of claims 29, 30, 32 or 33 or the vectors of the composition of claim 31 or 32 or obtainable by the method of any one of claims 1 to 27.

- 38. The transgenic plant cell of claim 37 which displays an increased cell division rate.
- 39. A transgenic plant or plant tissue comprising plant cells of claim 37 or 38 or obtainable by the method of any one of claims 1 to 27.
- 40. The transgenic plant of claim 39 which displays modified and/or accelerated and/or enhanced plant growth, root growth, shoot growth and/or yield or modified architecture compared to the corresponding wild type plant.
- 41. Harvestable parts or propagation material of a plant of claim 29 or 30 comprising the plant cell of claim 37 or 38 or the tissue of claim 39.
- 42. Use of at least one nucleic acid molecule or regulatory sequence as defined in any one of claims 1 to 21 or a nucleic acid molecule of claim 28, the vector of any one of claims 29, 30, 32 or 33 or the vectors of the composition of claim 31 or 32 for increasing cell division rates in plants, plant cells or plant tissue.
- 43. The use of claim 42, wherein said increased cell division rates result in increased or enhanced biomass, plant growth, root and/or shoot growth, seed setting, seed set, seed production, grain yield, yield of harvestable material, modified architecture, fruit size, nitrogen-fixing capability, nodule size, stem thickness, endosperm size, number of fruits per plant and/or initiation, promotion, stimulation or enhancement of seed development, tuber formation, shoot initiation, leaf initiation, inhibition of apical dominance and/or development.
- 44. A composition comprising nucleic acid molecules or regulatory sequences as defined in any one of claims 1 to 21, a nucleic acid molecule of claim 28, the vector of any one of claims 29, 30, 32 or 33,

the plant cells of claim 37 or 38, the tissue of claim 38 or the vectors of the composition of claim 31 or 32.

45. Use of at least one nucleic acid molecule or regulatory sequence as defined in any one of claims 1 to 21 or a nucleic acid molecule of claim 28, the vector of any one of claims 29, 30, 32 or 33 or the vectors of the composition of claims 31 or 32, the tissue of claim 39 or the plant cells of claim 37 or 38 for the production of more biomass, of secondary metabolites or additives for plant culturing in plant cell culture.

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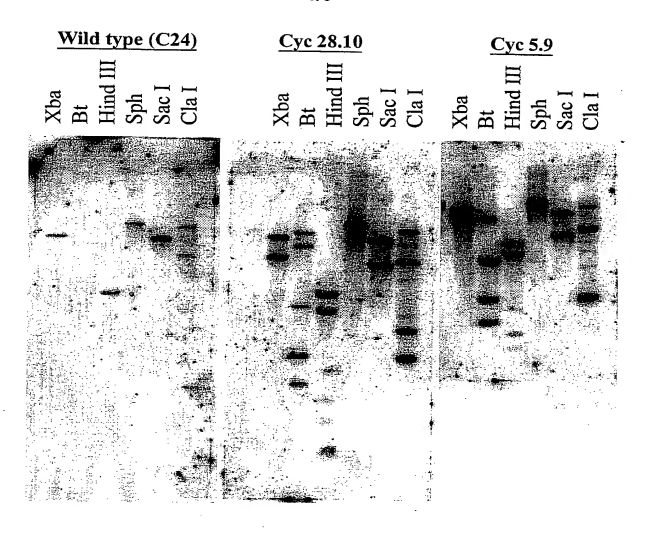


Fig. 1

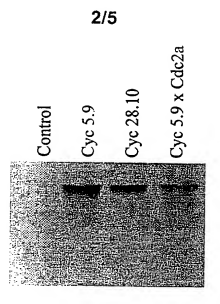


Fig. 2

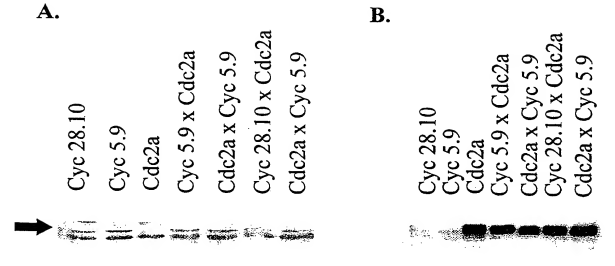


Fig. 3

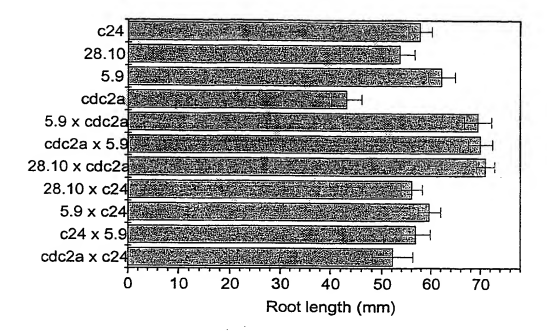


Fig. 4

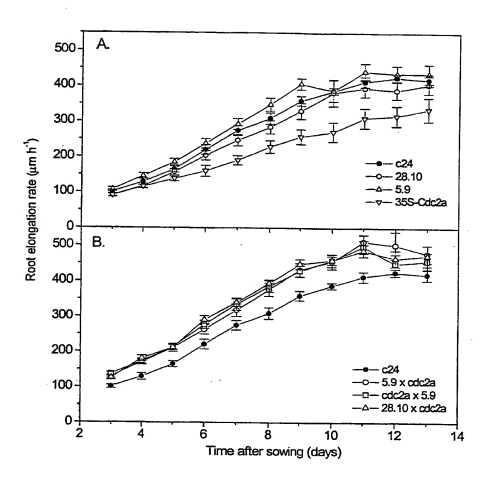


Fig. 5

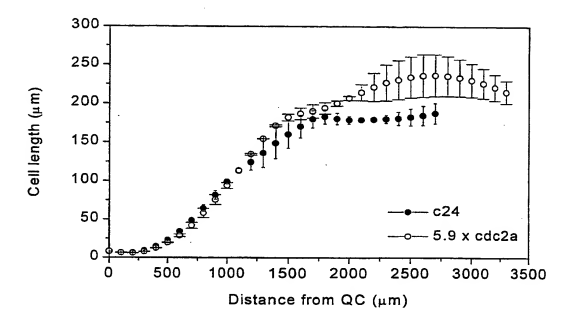


Fig. 6

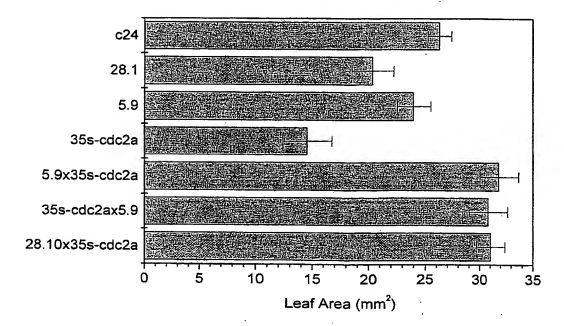


Fig. 7

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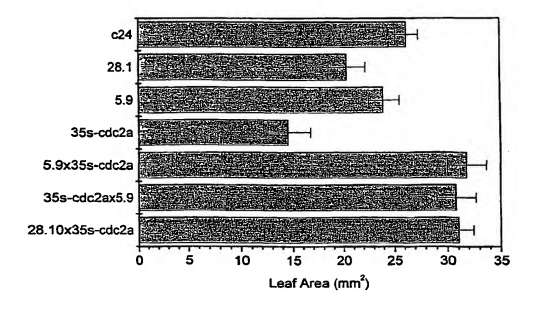
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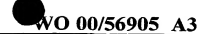
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(54) Title: METHOD FOR ENHANCING AND/OR IMPROVING PLANT GROWTH AND/OR YIELD OR MODIFYING PLANT ARCHITECTURE



(57) Abstract: Described is a method for promoting or modifying plant growth and/or yield and/or architecture in plants comprising the increased expression of at least two cell cycle interacting proteins, in particular of a protein kinase, e.g. CDK, and a cyclin. Transgenic plants are provided obtainable by this method and displaying increased cell division rates and growth rates. In addition, harvestable parts and propagation material of the above-mentioned plant as well as the use of the provided cells, tissues and plants for the production of biomass, secondary metabolites or additives for plant culturing in plant culture.

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Inte	al Application No
PC 1,	P 00/02441

A CLASSIFICATION OF SUBJECT MATTER
IPC 7 C12N15/29 C12N15/82 C12N9/12 C12N5/10 A01H5/00 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 7 C12N A01H Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal, BIOSIS C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. WO 97 02354 A (MEDICAL RES COUNCIL 28-36 ;LATHANGUE NICHOLAS BARRIE (GB)) 23 January 1997 (1997-01-23) the whole document A WO 98 41642 A (VEYLDER LIEVEN DE ; VLAAMS 1-45 INTERUNIV INST BIOTECH (BE); INZE DIRK () 24 September 1998 (1998-09-24) cited in the application see the whole document; esp. pp.8-16 WO 99 13083 A (VEYLDER LIEVEN DE ;INZE A 1-45 DIRK (BE); CROPDESIGN N V (BE); SEGERS GER) 18 March 1999 (1999-03-18) the whole document X Further documents are listed in the continuation of box C. Patent family members are listed in annex. Special categories of cited documents: "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention filing date cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention carnot be considered to involve an inventive step when the document is combined with one or more other such docu-"O" document referring to an oral disclosure, use, exhibition or other means ments, such combination being obvious to a person skilled in the art. document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 5 September 2000 21/09/2000 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Fax: (+31-70) 340-3016 Kania, T

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